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**C A** 

(54) Title: METHODS OF DIAGNOSIS AND TREATMENT OF ANDROGEN-DEPENDENT PROSTATE CANCER, PROSTATE CANCER UNDERGOING ANDROGEN-WITHDRAWAL, AND ANDROGEN-INDEPENDENT PROSTATE CANCER

(57) Abstract: Described herein are genes whose expression are up-regulated or down-regulated in prostate cancer. Also described are such genes whose expression is further up-regulated or down-regulated in drug-resistant prostate cancer cells. Related methods and compositions that can be used for diagnosis and treatment of prostate cancer are disclosed. Also described herein are methods that can be used to identify modulators of prostate cancer.

METHODS OF DIAGNOSIS AND TREATMENT OF ANDROGEN-DEPENDENT PROSTATE CANCER, PROSTATE CANCER UNDERGOING ANDROGEN WITHDRAWAL, AND ANDROGEN-INDEPENDENT PROSTATE CANCER

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# CROSS-REFERENCES TO RELATED APPLICATIONS

This application claims priority from the following applications: USSN 60/295,917, filed June 4, 2001, USSN 60/368,689, filed March 29, 2002; USSN 60/350,666, filed November 13, 2001; and USSN 60/372,246, filed April 12, 2002; each of which is incorporated herein by reference in its entirety.

#### FIELD OF THE INVENTION

The invention relates to the identification of nucleic acid and protein expression profiles and nucleic acids, products, and antibodies thereto that are involved in prostate cancer; and to the use of such expression profiles and compositions in the diagnosis, prognosis, and therapy of prostate cancer. The invention further relates to methods for identifying and using agents and/or targets that inhibit prostate cancer.

## BACKGROUND OF THE INVENTION

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Prostate cancer is the most frequently diagnosed cancer and the second leading cause of male cancer death in North America and northern Europe. Early detection of prostate cancer using a serum test for prostate-specific antigen (PSA) has dramatically improved the treatment of the disease (Oesterling (1992) J. Am. Med. Assoc. 267:2236-2238). Treatment of prostate cancer consists largely of surgical prostatectomy, radiation therapy, androgen ablation therapy and chemotherapy. Although many prostate cancer patients are effectively treated, the current therapies can all induce serious side effects which diminish quality of life. Patients who present with metastatic disease are most often treated with androgen-ablation therapy. Hormone blockade results in significant regression of the tumor. However, this treatment rarely cures the patient and invariably results in progression to androgen-

independent disease, which is incurable. Afrin and Stuart (1994) <u>J.S.C. Med. Assoc.</u> 90:231-236.

The identification of novel therapeutic targets and diagnostic markers is essential for improving the current treatment of prostate cancer patients. Recent advances in molecular medicine have increased the interest in tumor-specific cell surface antigens that could serve as targets for various immunotherapeutic or small molecule strategies. Antigens suitable for immunotherapeutic strategies should be highly expressed in cancer tissues and ideally not expressed in normal adult tissues. Expression in tissues that are dispensable for life, however, may be tolerated. Examples of such antigens include Her2/neu and the B-cell antigen CD20. Humanized monoclonal antibodies directed to Her2/neu (Herceptin) are currently in use for the treatment of metastatic breast cancer. Ross and Fletcher (1998) Stem Cells 16:413-428. Similarly, anti-CD20 monoclonal antibodies (Rituxin) are used to effectively treat non-Hodgkin's lymphoma. Maloney, et al. (1997) Blood 90:2188-2195; Leget and Czuczman (1998) Curr. Opin. Oncol. 10:548-551.

Several potential immunotherapeutic targets have been identified for prostate cancer. They include prostate-specific membrane antigen (PSMA) (Israeli, et al. (1993) Cancer Res. 53:227-230), prostate stem cell antigen (PSCA; Reiter, et al. (1998) Proc. Natl. Acad. Sci. USA 95:1735-1740), and serpentine transmembrane epithelial antigen of the prostate (STEAP; Hubert, et al. (1999) Proc. Natl. Acad. Sci. USA 96:14529-14534). PSMA is a type II transmembrane hydrolase with significant homology to a rat neuropeptidase (Carter, et al. (1996) Proc. Natl. Acad. Sci. USA 93:749-753). Antibodies directed towards PSMA are currently being used to detect metastasized prostate cancer as the Prostascint Scan (Sodee, et al. (1996) Clin. Nucl. Med. 21:759-767) and are also being evaluated for treatment of advanced disease (Gregorakis, et al. (1998) Semin. Urol. Oncol. 16:2-12; Liu, et al. (1998) Cancer Res. 58:4055-4060; Murphy, et al. (1998) J. Urol. 160:2396-2401). In a study on bone metastasis of prostate cancer, only 8 out of 18 patient samples expressed PSMA (Silver, et al. (1997) Clin. Cancer Res. 3:81-85). Therefore, it is clear that other targets need to be identified to manage metastasized disease. PSCA is a member of the Thy-1/Ly-6 family of glycosylphosphatidylinositol-linked plasma membrane proteins (Reiter, et al. (1998) Proc. Natl. Acad. Sci. USA 95:1735-1740). Immunohistochemical data shows that PSCA is upregulated in the majority of prostate cancer epithelia and is also detected in bone metastasis (Gu, et al. (2000) Oncogene 19:1288-1296). Recent work shows that antibodies directed to

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PSCA can prevent metastatic spread of prostate cancer in a mouse model (Saffran, et al. (2001) Proc. Natl. Acad. Sci. USA 98:2658-2663). STEAP is a multi-transmembrane prostate-specific protein that may function as a channel or transporter protein (Hubert, et al. (1999) Proc. Natl. Acad. Sci. USA 96:14529-14534). Its protein expression is specific to the basolateral membranes of normal prostate and prostate cancer epithelia. STEAP expression was most highly concentrated at cell-cell boundaries, implying a potential function in intercellular communication. Therapeutic monoclonal antibodies have so far not been reported for STEAP.

### SUMMARY OF THE INVENTION

The present invention therefore provides nucleotide sequences of genes that are upand down-regulated in androgen-independent prostate cancer cells or prostate cells
undergoing androgen withdrawal. Such genes are useful for diagnostic purposes, and also as
targets for screening for therapeutic compounds that modulate prostate cancer, such as
hormones or antibodies. Other aspects of the invention will become apparent to the skilled
artisan by the following description of the invention.

In one aspect, the present invention provides a method of detecting an androgen independent prostate cancer-associated transcript in a cell from a patient, the method comprising contacting a biological sample from the patient with a polynucleotide that selectively hybridizes to nucleic acid molecule comprising a sequence at least 80% identical to a sequence as shown in Tables 1A-4.

In one embodiment, the present invention provides a method of determining the level of a prostate cancer associated transcript in a cell from a patient.

In one embodiment, the present invention provides a method of detecting a prostate cancer-associated transcript in a cell from a patient, the method comprising contacting a biological sample from the patient with a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1A-4.

In various embodiments, the polynucleotide selectively hybridizes to a sequence at least 95% identical to a sequence as shown in Tables 1A-4; the polynucleotide comprises a sequence as shown in Tables 1A-4; the biological sample is a tissue sample; the biological sample comprises isolated nucleic acids, e.g., mRNA; the polynucleotide is labeled, e.g., with a fluorescent label; the polynucleotide is immobilized on a solid surface; the patient is

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undergoing a therapeutic regimen to treat prostate cancer; the patient is suspected of having metastatic prostate cancer; the patient is a human; the patient is suspected of having a taxol-resistant cancer; or the prostate cancer associated transcript is mRNA.

In other embodiments, the method further comprises the step of amplifying nucleic acids before the step of contacting the biological sample with the polynucleotide.

In another aspect, the present invention provides a method of monitoring the efficacy of a therapeutic treatment of prostate cancer, the method comprising the steps of: (i) providing a biological sample from a patient undergoing the therapeutic treatment; and (ii) determining the level of a prostate cancer-associated transcript in the biological sample by contacting the biological sample with a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1A-4, thereby monitoring the efficacy of the therapy. In a further embodiment, the patient has metastatic prostate cancer. In a further embodiment, the patient has a drug resistant (e.g., taxol resistant) form of prostate cancer.

In one embodiment, the method further comprises the step of: (iii) comparing the level of the prostate cancer-associated transcript to a level of the prostate cancer-associated transcript in a biological sample from the patient prior to, or earlier in, the therapeutic treatment.

Additionally, provided herein is a method of evaluating the effect of a candidate prostate cancer drug comprising administering the drug to a patient and removing a cell sample from the patient. The expression profile of the cell is then determined. This method may further comprise comparing the expression profile to an expression profile of a healthy individual. In a preferred embodiment, said expression profile includes a gene of Tables 1A-4.

In one aspect, the present invention provides an isolated nucleic acid molecule consisting of a polynucleotide sequence as shown in Tables 1A-4.

In one embodiment, an expression vector or cell comprises the isolated nucleic acid.

In one aspect, the present invention provides an isolated polypeptide which is encoded by a nucleic acid molecule having polynucleotide sequence as shown in Tables 1A-4.

In another aspect, the present invention provides an antibody that specifically binds to an isolated polypeptide which is encoded by a nucleic acid molecule having polynucleotide sequence as shown in Tables 1A-4.

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In certain embodiments, the antibody is conjugated to an effector component, e.g., a fluorescent label, a radioisotope or a cytotoxic chemical; the antibody is an antibody fragment; or the antibody is humanized.

In one aspect, the present invention provides a method of detecting a prostate cancer cell in a biological sample from a patient, the method comprising contacting the biological sample with an antibody as described herein.

In another aspect, the present invention provides a method of detecting antibodies specific to prostate cancer in a patient, the method comprising contacting a biological sample from the patient with a polypeptide encoded by a nucleic acid comprising a sequence from Tables 1A-4.

In another aspect, the present invention provides a method for identifying a compound that modulates a prostate cancer-associated polypeptide, the method comprising the steps of:
a) contacting the compound with a prostate cancer-associated polypeptide, the polypeptide encoded by a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1A-4; and b) determining the functional effect of the compound upon the polypeptide.

In one embodiment, the functional effect is a physical effect, an enzymatic effect, or a chemical effect.

In one embodiment, the polypeptide is expressed in a eukaryotic host cell or cell membrane. In another embodiment, the polypeptide is recombinant.

In one embodiment, the functional effect is determined by measuring ligand binding to the polypeptide.

In another aspect, the present invention provides a method of inhibiting proliferation of a prostate cancer-associated cell to treat prostate cancer in a patient, the method comprising the step of administering to the subject a therapeutically effective amount of a compound identified as described herein.

In one embodiment, the compound is an antibody.

In another aspect, the present invention provides a drug screening assay comprising the steps of: a) administering a test compound to a mammal having prostate cancer or to a cell sample isolated therefrom; b) comparing the level of gene expression of a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1A-4 in a treated cell or mammal with the level of gene expression of the

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polynucleotide in a control cell sample or mammal, wherein a test compound that modulates the level of expression of the polynucleotide is a candidate for the treatment of prostate cancer.

In one embodiment, the control is a mammal with prostate cancer or a cell sample therefrom that has not been treated with the test compound. In another embodiment, the control is a normal cell or mammal.

In one embodiment, the test compound is administered in varying amounts or concentrations. In another embodiment, the test compound is administered for varying time periods. In another embodiment, the comparison can occur after addition or removal of the drug candidate.

In one embodiment, the levels of a plurality of polynucleotides that selectively hybridize to a sequence at least 80% identical to a sequence as shown in Tables 1A-4 are individually compared to their respective levels in a control cell sample or mammal. In a preferred embodiment the plurality of polynucleotides is from three to ten.

In another aspect, the present invention provides a method for treating a mammal having prostate cancer comprising administering a compound identified by the assay described herein.

In another aspect, the present invention provides a pharmaceutical composition for treating a mammal having prostate cancer, the composition comprising a compound identified by the assay described herein and a physiologically acceptable excipient.

In one aspect, the present invention provides a method of screening drug candidates by providing a cell expressing a gene that is up- and down-regulated as in a prostate cancer. In one embodiment, a gene is selected from Tables 1A-4. The method further includes adding a drug candidate to the cell and determining the effect of the drug candidate on the expression of the expression profile gene.

In one embodiment, the method of screening drug candidates includes comparing the level of expression in the absence of the drug candidate to the level of expression in the presence of the drug candidate, wherein the concentration of the drug candidate can vary when present, and wherein the comparison can occur after addition or removal of the drug candidate. In a preferred embodiment, the cell expresses at least two expression profile genes. The profile genes may show an increase or decrease.

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Also provided is a method of evaluating the effect of a candidate prostate cancer drug comprising administering the drug to a transgenic animal expressing or over-expressing the prostate cancer modulatory protein, or an animal lacking the prostate cancer modulatory protein, for example as a result of a gene knockout.

Moreover, provided herein is a biochip comprising one or more nucleic acid segments of Tables 1A-4, wherein the biochip comprises fewer than 1000 nucleic acid probes. Preferably, at least two nucleic acid segments are included. More preferably, at least three nucleic acid segments are included.

Furthermore, a method of diagnosing a disorder associated with prostate cancer is provided. The method comprises determining the expression of a gene of Tables 1A-4, in a first tissue type of a first individual, and comparing the distribution to the expression of the gene from a second normal tissue type from the first individual or a second unaffected individual. A difference in the expression indicates that the first individual has a disorder associated with prostate cancer.

In a further embodiment, the biochip also includes a polynucleotide sequence of a gene that is not up- and down-regulated in prostate cancer.

In one embodiment a method for screening for a bioactive agent capable of interfering with the binding of a prostate cancer modulating protein (prostate cancer modulatory protein) or a fragment thereof and an antibody which binds to said prostate cancer modulatory protein or fragment thereof. In a preferred embodiment, the method comprises combining a prostate cancer modulatory protein or fragment thereof, a candidate bioactive agent and an antibody which binds to said prostate cancer modulatory protein or fragment thereof. The method further includes determining the binding of said prostate cancer modulatory protein or fragment thereof and said antibody. Wherein there is a change in binding, an agent is identified as an interfering agent. The interfering agent can be an agonist or an antagonist. Preferably, the agent inhibits prostate cancer.

Also provided herein are methods of eliciting an immune response in an individual. In one embodiment a method provided herein comprises administering to an individual a composition comprising a prostate cancer modulating protein, or a fragment thereof. In another embodiment, the protein is encoded by a nucleic acid selected from those of Tables 1A-4.

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Further provided herein are compositions capable of eliciting an immune response in an individual. In one embodiment, a composition provided herein comprises a prostate cancer modulating protein, preferably encoded by a nucleic acid of Tables 1A-4, or a fragment thereof, and a pharmaceutically acceptable carrier. In another embodiment, said composition comprises a nucleic acid comprising a sequence encoding a prostate cancer modulating protein, preferably selected from the nucleic acids of Tables 1A-4 and a pharmaceutically acceptable carrier.

Also provided are methods of neutralizing the effect of a prostate cancer protein, or a fragment thereof, comprising contacting an agent specific for said protein with said protein in an amount sufficient to effect neutralization. In another embodiment, the protein is encoded by a nucleic acid selected from those of Tables 1A-4. In another aspect of the invention, a method of treating an individual for prostate cancer is provided. In one embodiment, the method comprises administering to said individual an inhibitor of a prostate cancer modulating protein. In another embodiment, the method comprises administering to a patient having prostate cancer an antibody to a prostate cancer modulating protein conjugated to a therapeutic moiety. Such a therapeutic moiety can be a cytotoxic agent or a radioisotope.

### DETAILED DESCRIPTION OF THE INVENTION

In accordance with the objects outlined above, the present invention provides novel methods for diagnosis and evaluation of androgen-dependent prostate cells (malignant or non-malignant), prostate cells undergoing androgen withdrawal, and androgen-independent prostate cancer, as well as methods for treating androgen-dependent prostate cells (malignant or non-malignant), prostate cancer undergoing androgen withdrawal, and androgen-independent prostate cancer. The current Specification incorporates the text of USSN 09/976,858, filed October 12, 2001, USSN 60/295,917, filed June 4, 2001, USSN 60/368,689, filed March 29, 2002; USSN 60/350,666, filed November 13, 2001; and USSN 60/372,246, filed April 12, 2002.

Table 1A provides unigene cluster identification numbers for the nucleotide sequence of genes that exhibit increased or decreased expression in androgen-independent prostate cancer samples. Table 1A also provides an exemplar accession number that provides a nucleotide sequence that is part of the unigene cluster. The expression patterns of the genes of Table 1A can be broadly defined into the following categories:

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Genes that are expressed early in the time course, then drop off in expression, and then express again with emergence of androgen-independence (hi-lo-hi pattern in table 1A). Genes that are expressed early in the time course, then drop off in expression, and do not express again with emergence of androgen-independence (hi-lo-lo pattern in 1A). Genes that are not expressed early in the time course, but express only with emergence of androgen-independence (lo-lo-hi pattern in table 1A). Genes that are not expressed early in the time course, but then express as androgen is withdrawn and continue to express with emergence of androgen-independence (lo-hi-hi pattern in table 1A). Genes that are not expressed early in the time course, but then express as androgen is withdrawn and drop off again with emergence of androgen-independence (lo-hi-lo pattern in table 1A).

Tables 2A-C provide unigene cluster identification numbers for the nucleotide sequence of genes that exhibit increased or decreased expression in androgen-dependent prostate cancer, prostate cancer undergoing androgen withdrawal and androgen-independent prostate cancer. Tables 2A-C also provide an exemplar accession number that provides a nucleotide sequence that is part of the unigene cluster. The expression patterns of the genes of Tables 2A-C can be broadly defined into the following 6 categories:

Genes that are expressed early in the time course of androgen withdrawal, then drop off in expression, and then express again with emergence of androgen-independence (hi-lo-lo-hi pattern in Table 2A). Genes that are expressed early in the time course, then drop off in expression immediately after androgen-withdrawal, and do not express again with emergence of androgen-independence (hi-lo-lo-lo pattern in Table 2A). Genes that are expressed early in the time course, then drop off in expression after several days of androgen withdrawal, and do not express again with emergence of androgen-independence (hi-hi-lo-lo pattern in Table 2A). Genes that are not expressed early in the time course, but express only with emergence of androgen-independence (lo-lo-hi pattern in Table 2A). Genes that are not expressed early in the time course, but then express as androgen is withdrawn and continue to express with emergence of androgen-independence (lo-lo-hi-hi pattern in Table 2A). Genes that are not expressed early in the time course, but then express as androgen is withdrawn and drop off again with emergence of androgen-independence (lo-lo-hi-hi pattern in Table 2A).

**Definitions** 

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The term "androgen ablation therapy" refers to techniques for the removal or destruction of sources of male hormones, such as testosterone. These techniques include, for example, 1) surgical removal of the testicles, 2) medications such as gonadatropin releasing hormone analogs that inhibit testosterone production, or 3) anti-androgenic drugs that block androgen receptors.

The term "androgen-independent prostate cancer protein" or "androgen-independent prostate cancer polynucleotide" or "androgen-independent prostate cancer-associated transcript" refers to nucleic acid and polypeptide polymorphic variants, alleles, mutants, and interspecies homologues that: (1) have a nucleotide sequence that has greater than about 60% nucleotide sequence identity, 65%, 70%, 75%, 80%, 85%, 90%, preferably 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% or greater nucleotide sequence identity, preferably over a region of over a region of at least about 25, 50, 100, 200, 500, 1000, or more nucleotides, to a nucleotide sequence of or associated with a unigene cluster of Tables 1A-4; (2) bind to antibodies, e.g., polyclonal antibodies, raised against an immunogen comprising an amino acid sequence encoded by a nucleotide sequence of or associated with a unigene cluster of Tables 1A-4 and conservatively modified variants thereof; (3) specifically hybridize under stringent hybridization conditions to a nucleic acid sequence, or the complement thereof of Tables 1A-4 and conservatively modified variants thereof; or (4) have an amino acid sequence that has greater than about 60% amino acid sequence identity, 65%, 70%, 75%, 80%, 85%, 90%, preferably 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% or greater amino sequence identity, preferably over a region of over a region of at least about 25, 50, 100, 200, 500, 1000, or more amino acid, to an amino acid sequence encoded by a nucleotide sequence of or associated with a unigene cluster of Tables 1A-4. These polynucleotides or proteins may also be expressed during a period following androgen withdrawal. A polynucleotide or polypeptide sequence is typically from a mammal including, but not limited to, primate, e.g., human; rodent, e.g., rat, mouse, hamster; cow, pig, horse, sheep, or other mammal. A "prostate cancer polypeptide" and a "prostate cancer polynucleotide," include both naturally occurring or recombinant forms, and may refer to those polypeptides or polynucleotides which are expressed in prostate proliferative cells.

A "full length" prostate cancer protein or nucleic acid refers to a prostate cancer polypeptide or polynucleotide sequence, or a variant thereof, that contains the elements normally contained in one or more naturally occurring, wild type prostate cancer

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polynucleotide or polypeptide sequences. The "full length" may be prior to, or after, various stages of post-translation processing or splicing, including alternative splicing.

"Biological sample" as used herein is a sample of biological tissue or fluid that contains nucleic acids or polypeptides, e.g., of a prostate cancer protein, polynucleotide or transcript. Such samples include, but are not limited to, tissue isolated from primates, e.g., humans, or rodents, e.g., mice, and rats. Biological samples may also include sections of tissues such as biopsy and autopsy samples, frozen sections taken for histology purposes, blood, plasma, serum, sputum, stool, tears, mucus, hair, skin, etc. Biological samples also include explants and primary and/or transformed cell cultures derived from patient tissues. A biological sample is typically obtained from a eukaryotic organism, most preferably a mammal such as a primate e.g., chimpanzee or human; cow; dog; cat; a rodent, e.g., guinea pig, rat, mouse; rabbit; or a bird; reptile; or fish.

"Providing a biological sample" means to obtain a biological sample for use in methods described in this invention. Most often, this will be done by removing a sample of cells from an animal, but can also be accomplished by using previously isolated cells (e.g., isolated by another person, at another time, and/or for another purpose), by collecting a sample which contains a soluble polypeptide or nucleic acid derived from a prostate cell, or by performing the methods of the invention in vivo. Archival tissues, having treatment or outcome history, will be particularly useful.

The terms "identical" or percent "identity," in the context of two or more nucleic acids or polypeptide sequences, refer to two or more sequences or subsequences that are the same or have a specified percentage of amino acid residues or nucleotides that are the same (i.e., about 60% identity, preferably 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or higher identity over a specified region, when compared and aligned for maximum correspondence over a comparison window or designated region) as measured using a BLAST or BLAST 2.0 sequence comparison algorithms with default parameters described below, or by manual alignment and visual inspection (see, e.g., NCBI web site http://www.ncbi.nlm.nih.gov/BLAST/ or the like). Such sequences are then said to be "substantially identical." This definition also refers to, or may be applied to, the compliment of a test sequence. The definition also includes sequences that have deletions and/or additions, as well as those that have substitutions, as well as naturally occurring, e.g., polymorphic or allelic variants, and man-made variants. As described below, the preferred

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algorithms can account for gaps and the like. Preferably, identity exists over a region that is at least about 25 amino acids or nucleotides in length, or more preferably over a region that is 50-100 amino acids or nucleotides in length.

For sequence comparison, typically one sequence acts as a reference sequence, to which test sequences are compared. When using a sequence comparison algorithm, test and reference sequences are entered into a computer, subsequence coordinates are designated, if necessary, and sequence algorithm program parameters are designated. Preferably, default program parameters can be used, or alternative parameters can be designated. The sequence comparison algorithm then calculates the percent sequence identities for the test sequences relative to the reference sequence, based on the program parameters.

A "comparison window", as used herein, includes reference to a segment of one of the number of contiguous positions selected from the group consisting typically of from 20 to 600, usually about 50 to about 200, more usually about 100 to about 150 in which a sequence may be compared to a reference sequence of the same number of contiguous positions after the two sequences are optimally aligned. Methods of alignment of sequences for comparison are well-known in the art. Optimal alignment of sequences for comparison can be conducted, e.g., by the local homology algorithm of Smith and Waterman (1981) Appl. Math. 2:482, by the homology alignment algorithm of Needleman and Wunsch (1970) J. Mol. Biol. 48:443-453, by the search for similarity method of Pearson and Lipman (1988) Proc. Nat'l. Acad. Sci. USA 85:2444-2448, by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI), or by manual alignment and visual inspection (see, e.g., Ausubel, et al. (eds. 1995 and supplements) Current Protocols in Molecular Biology Lippincott).

Preferred examples of algorithms that are suitable for determining percent sequence identity and sequence similarity include the BLAST and BLAST 2.0 algorithms, which are described in Altschul, et al. (1977) Nuc. Acids Res. 25:3389-3402 and Altschul, et al. (1990) J. Mol. Biol. 215:403-410. BLAST and BLAST 2.0 are used, with the parameters described herein, to determine percent sequence identity for the nucleic acids and proteins of the invention. Software for performing BLAST analyses is publicly available through the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/). This algorithm involves first identifying high scoring sequence pairs (HSPs) by identifying short

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words of length W in the query sequence, which either match or satisfy some positive-valued threshold score T when aligned with a word of the same length in a database sequence. T is referred to as the neighborhood word score threshold (Altschul, et al., supra). These initial neighborhood word hits act as seeds for initiating searches to find longer HSPs containing them. The word hits are extended in both directions along each sequence for as far as the cumulative alignment score can be increased. Cumulative scores are calculated using, e.g., for nucleotide sequences, the parameters M (reward score for a pair of matching residues; always > 0) and N (penalty score for mismatching residues; always < 0). For amino acid sequences, a scoring matrix is used to calculate the cumulative score. Extension of the word hits in each direction are halted when: the cumulative alignment score falls off by the quantity X from its maximum achieved value; the cumulative score goes to zero or below, due to the accumulation of one or more negative-scoring residue alignments; or the end of either sequence is reached. The BLAST algorithm parameters W, T, and X determine the sensitivity and speed of the alignment. The BLASTN program (for nucleotide sequences) uses as defaults a wordlength (W) of 11, an expectation (E) of 10, M=5, N=-4 and a comparison of both strands. For amino acid sequences, the BLASTP program uses as defaults a wordlength of 3, and expectation (E) of 10, and the BLOSUM62 scoring matrix (see Henikoff and Henikoff (1989) Proc. Natl. Acad. Sci. USA 89:10915-919) alignments (B) of 50, expectation (E) of 10, M=5, N=-4, and a comparison of both strands.

The BLAST algorithm also performs a statistical analysis of the similarity between two sequences (see, e.g., Karlin and Altschul (1993) Proc. Nat'l. Acad. Sci. USA 90:5873-5787). One measure of similarity provided by the BLAST algorithm is the smallest sum probability (P(N)), which provides an indication of the probability by which a match between two nucleotide or amino acid sequences would occur by chance. For example, a nucleic acid is considered similar to a reference sequence if the smallest sum probability in a comparison of the test nucleic acid to the reference nucleic acid is less than about 0.2, more preferably less than about 0.01, and most preferably less than about 0.001. Log values may be large negative numbers, e.g., 5, 10, 20, 30, 40, 40, 70, 90, 110, 150, 170, etc.

An indication that two nucleic acid sequences or polypeptides are substantially identical is that the polypeptide encoded by the first nucleic acid is immunologically cross reactive with the antibodies raised against the polypeptide encoded by the second nucleic acid, as described below. Thus, a polypeptide is typically substantially identical to a second

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polypeptide, e.g., where the two peptides differ only by conservative substitutions. Another indication that two nucleic acid sequences are substantially identical is that the two molecules or their complements hybridize to each other under stringent conditions, as described below. Yet another indication that two nucleic acid sequences are substantially identical is that the same primers can be used to amplify the sequences.

A "host cell" is a naturally occurring cell or a transformed cell that contains an expression vector and supports the replication or expression of the expression vector. Host cells may be cultured cells, explants, cells in vivo, and the like. Host cells may be prokaryotic cells such as E. coli, or eukaryotic cells such as yeast, insect, amphibian, or mammalian cells such as CHO, HeLa, and the like (see, e.g., the American Type Culture Collection catalog or web site, www.atcc.org).

The terms "isolated," "purified," or "biologically pure" refer to material that is substantially or essentially free from components that normally accompany it as found in its native state. Purity and homogeneity are typically determined using analytical chemistry techniques such as polyacrylamide gel electrophoresis or high performance liquid chromatography. A protein or nucleic acid that is the predominant species present in a preparation is substantially purified. In particular, an isolated nucleic acid is separated from some open reading frames that naturally flank the gene and encode proteins other than protein encoded by the gene. The term "purified" in some embodiments denotes that a nucleic acid or protein gives rise to essentially one band in an electrophoretic gel. Preferably, it means that the nucleic acid or protein is at least 85% pure, more preferably at least 95% pure, and most preferably at least 99% pure. "Purify" or "purification" in other embodiments means removing at least one contaminant from the composition to be purified. In this sense, purification does not require that the purified compound be homogenous, e.g., 100% pure.

The terms "polypeptide," "peptide," and "protein" are used interchangeably herein to refer to a polymer of amino acid residues. The terms apply to amino acid polymers in which one or more amino acid residue is an artificial chemical mimetic of a corresponding naturally occurring amino acid, as well as to naturally occurring amino acid polymers, those containing modified residues, and non-naturally occurring amino acid polymer. Certain diagnostic methods may evaluate secreted or breakdown products present only because the producing cell is present, and would otherwise be absent in a normal individual.

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The term "amino acid" refers to naturally occurring and synthetic amino acids, as well as amino acid analogs and amino acid mimetics that function similarly to the naturally occurring amino acids. Naturally occurring amino acids are those encoded by the genetic code, as well as those amino acids that are later modified, e.g., hydroxyproline, γ-carboxyglutamate, and O-phosphoserine. Amino acid analogs refers to compounds that have the same basic chemical structure as a naturally occurring amino acid, e.g., an α carbon that is bound to a hydrogen, a carboxyl group, an amino group, and an R group, e.g., homoserine, norleucine, methionine sulfoxide, methionine methyl sulfonium. Such analogs may have modified R groups (e.g., norleucine) or modified peptide backbones, but retain the same basic chemical structure as a naturally occurring amino acid. Amino acid mimetics refers to chemical compounds that have a structure that is different from the general chemical structure of an amino acid, but that functions similarly to a naturally occurring amino acid.

Amino acids may be referred to herein by either their commonly known three letter symbols or by the one-letter symbols recommended by the IUPAC-IUB Biochemical Nomenclature Commission. Nucleotides, likewise, may be referred to by their commonly accepted single-letter codes.

"Conservatively modified variants" applies to both amino acid and nucleic acid sequences. With respect to particular nucleic acid sequences, conservatively modified variants refers to those nucleic acids which encode identical or essentially identical amino acid sequences, or where the nucleic acid does not encode an amino acid sequence, to essentially identical or associated, e.g., naturally contiguous, sequences. Because of the degeneracy of the genetic code, a large number of functionally identical nucleic acids encode most proteins. For instance, the codons GCA, GCC, GCG, and GCU encode the amino acid alanine. Thus, at every position where an alanine is specified by a codon, the codon can be altered to another of the corresponding codons described without altering the encoded polypeptide. Such nucleic acid variations are "silent variations," which are one species of conservatively modified variations. Every nucleic acid sequence herein which encodes a polypeptide also describes silent variations of the nucleic acid. One of skill will recognize that in certain contexts each codon in a nucleic acid (except AUG, which is ordinarily the only codon for tryptophan) can be modified to yield a functionally identical molecule. Accordingly, often silent variations of

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a nucleic acid which encodes a polypeptide is implicit in a described sequence with respect to the expression product, but not with respect to actual probe sequences.

As to amino acid sequences, one of skill will recognize that individual substitutions, deletions or additions to a nucleic acid, peptide, polypeptide, or protein sequence which alters, adds or deletes a single amino acid or a small percentage of amino acids in the encoded sequence is a "conservatively modified variant" where the alteration results in the substitution of an amino acid with a chemically similar amino acid. Conservative substitutions providing functionally similar amino acids are well known in the art. Such conservatively modified variants are in addition to and do not exclude polymorphic variants, interspecies homologs, and alleles of the invention, typically conservative substitutions for one another: 1) Alanine (A), Glycine (G); 2) Aspartic acid (D), Glutamic acid (E); 3) Asparagine (N), Glutamine (Q); 4) Arginine (R), Lysine (K); 5) Isoleucine (I), Leucine (L), Methionine (M), Valine (V); 6) Phenylalanine (F), Tyrosine (Y), Tryptophan (W); 7) Serine (S), Threonine (T); and 8) Cysteine (C), Methionine (M) (see, e.g., Creighton (1984) Proteins Freeman).

Macromolecular structures such as polypeptide structures can be described in terms of various levels of organization. For a general discussion of this organization, see, e.g., Alberts, et al. (2001) Molecular Biology of the Cell (4th ed.) and Cantor and Schimmel (1980) Biophysical Chemistry Part I: The Conformation of Biological Macromolecules Freeman. "Primary structure" refers to the amino acid sequence of a particular peptide. "Secondary structure" refers to locally ordered, three dimensional structures within a polypeptide. These structures are commonly known as domains. Domains are portions of a polypeptide that often form a compact unit of the polypeptide and are typically 25 to approximately 500 amino acids long. Typical domains are made up of sections of lesser organization such as stretches of  $\beta$ -sheet and  $\alpha$ -helices. "Tertiary structure" refers to the complete three dimensional structure of a polypeptide monomer. "Quaternary structure" refers to the three dimensional structure formed, usually by the noncovalent association of independent tertiary units. Anisotropic terms are also known as energy terms.

"Nucleic acid" or "oligonucleotide" or "polynucleotide" or grammatical equivalents used herein means at least two nucleotides covalently linked together. Oligonucleotides are typically from about 5, 6, 7, 8, 9, 10, 12, 15, 25, 30, 40, 50 or more nucleotides in length, up to about 100 nucleotides in length. Nucleic acids and polynucleotides are a polymers of virtually any length, including longer lengths, e.g., 200, 300, 500, 1000, 2000, 3000, 5000,

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7000, 10,000, etc. A nucleic acid of the present invention will generally contain phosphodiester bonds, although in some cases, nucleic acid analogs are included that may have alternate backbones, comprising, e.g., phosphoramidate, phosphorothioate, phosphorodithioate, or O-methylphosphoroamidite linkages (see Eckstein (1992) Oligonucleotides and Analogues: A Practical Approach, Oxford University Press); and peptide nucleic acid backbones and linkages. Other analog nucleic acids include those with positive backbones; non-ionic backbones, and non-ribose backbones, including those described in U.S. Patent Nos. 5,235,033 and 5,034,506, and Chapters 6 and 7, ASC Symposium Series 580, Sanghvi and Cook (eds. 1994) Carbohydrate Modifications in Antisense Research ACS Symposium Series 580. Nucleic acids containing one or more carbocyclic sugars are also included within one definition of nucleic acids. Modifications of the ribose-phosphate backbone may be done for a variety of reasons, e.g., to increase the stability and half-life of such molecules in physiological environments or as probes on a biochip. Mixtures of naturally occurring nucleic acids and analogs can be made; alternatively, mixtures of different nucleic acid analogs, and mixtures of naturally occurring nucleic acids and analogs may be made.

A variety of references disclose such nucleic acid analogs, including, for example, phosphoramidate (Beaucage, et al. (1993) Tetrahedron 49(10):1925-1963 and references therein; Letsinger (1970) J. Org. Chem. 35:3800-3803; Sprinzl, et al. (1977) Eur. J. Biochem. 81:579-589; Letsinger, et al. (1986) Nucl. Acids Res. 14:3487-499; Sawai, et al (1984) 20 Chem. Lett. 805, Letsinger, et al. (1988) J. Am. Chem. Soc. 110:4470-4471; and Pauwels, et al. (1986) Chemica Scripta 26:141-149), phosphorothioate (Mag, et al. (1991) Nucleic Acids Res. 19:1437-441; and U.S. Patent No. 5,644,048), phosphorodithioate (Briu, et al. (1989) J. Am. Chem. Soc. 111:2321-xxx, O-methylphosphoroamidite linkages (see Eckstein (1992) 25 Oligonucleotides and Analogues: A Practical Approach Oxford University Press), and peptide nucleic acid backbones and linkages (see Egholm (1992) J. Am. Chem. Soc. 114:1895-1897; Meier, et al. (1992) Chem. Int. Ed. Engl. 31:1008-1010; Nielsen (1993) Nature 365:566-568; Carlsson, et al. (1996) Nature 380:207, each of which is incorporated by reference). Other analog nucleic acids include those with positive backbones (Denpcy, et al. (1995) Proc. Natl. Acad. Sci. USA 92:6097-101; non-ionic backbones (U.S. Patent Nos. 30 5,386,023, 5,637,684, 5,602,240, 5,216,141 and 4,469,863; Kiedrowshi, et al. (1991) Angew. Chem. Intl. Ed. English 30:423-426; Letsinger, et al. (1988) J. Am. Chem. Soc. 110:4470;

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Letsinger, et al. (1994) Nucleoside and Nucleotide 13:1597-xxx; Chapters 2 and 3 in Sanghvi and Cook (eds. 1994) Carbohydrate Modifications in Antisense Research ACS Symposium Series 580; Mesmaeker, et al. (1994) Bioorganic and Medicinal Chem. Lett. 4:395-xxx; Jeffs, et al. (1994) J. Biomolecular NMR 34:17; Horn (1996) Tetrahedron Lett. 37:743-xxx) and non-ribose backbones, including those described in U.S. Patent Nos. 5,235,033 and 5,034,506, and Chapters 6 and 7 in Sanghvi and Cook (eds. 1994) Carbohydrate Modifications in Antisense Research ACS Symposium Series 580. Nucleic acids containing one or more carbocyclic sugars are also included within one definition of nucleic acids (see Jenkins, et al. (1995) Chem. Soc. Rev. xx:169-176). Several nucleic acid analogs are described in Rawls (p. 35, June 2, 1997) C&E News. Each of these references is hereby expressly incorporated by reference.

Particularly preferred are peptide nucleic acids (PNA) which includes peptide nucleic acid analogs. These backbones are substantially non-ionic under neutral conditions, in contrast to the highly charged phosphodiester backbone of naturally occurring nucleic acids. This results in two advantages. First, the PNA backbone exhibits improved hybridization kinetics. PNAs have larger changes in the melting temperature (T<sub>m</sub>) for mismatched versus perfectly matched base pairs. DNA and RNA typically exhibit a 2-4° C drop in T<sub>m</sub> for an internal mismatch. With the non-ionic PNA backbone, the drop is closer to 7-9° C. Similarly, due to their non-ionic nature, hybridization of the bases attached to these backbones is relatively insensitive to salt concentration. In addition, PNAs are not degraded by cellular enzymes, and thus can be more stable.

The nucleic acids may be single stranded or double stranded, as specified, or contain portions of both double stranded or single stranded sequence. As will be appreciated by those in the art, the depiction of a single strand also defines the sequence of the complementary strand; thus the sequences described herein also provide the complement of the sequence. The nucleic acid may be DNA, both genomic and cDNA, RNA or a hybrid, where the nucleic acid may contain combinations of deoxyribo- and ribo-nucleotides, and combinations of bases, including uracil, adenine, thymine, cytosine, guanine, inosine, xanthine hypoxanthine, isocytosine, isoguanine, etc. "Transcript" typically refers to a naturally occurring RNA, e.g., a pre-mRNA, hnRNA, or mRNA. As used herein, the term "nucleoside" includes nucleotides and nucleoside and nucleotide analogs, and modified nucleosides such as amino modified nucleosides. In addition, "nucleoside" includes non-naturally occurring analog structures.

Thus, e.g., the individual units of a peptide nucleic acid, each containing a base, are referred to herein as a nucleoside.

A "label" or a "detectable moiety" is a composition detectable by spectroscopic, photochemical, biochemical, immunochemical, chemical, or other physical means. For example, useful labels include <sup>32</sup>P, fluorescent dyes, electron-dense reagents, enzymes (e.g., as commonly used in an ELISA), biotin, digoxigenin, or haptens and proteins or other entities which can be made detectable, e.g., by incorporating a radiolabel into the peptide or used to detect antibodies specifically reactive with the peptide. The labels may be incorporated into the prostate cancer nucleic acids, proteins, and antibodies at virtually any position. Many methods for conjugating the antibody to the label may be employed, including those methods described by Hunter, et al. (1962) Nature, 144:945; David, et al. (1974) Biochemistry 13:1014-1021; Pain, et al. (1981) J. Immunol. Meth. 40:219-230; and Nygren (1982) J. Histochem. and Cytochem. 30:407-412.

An "effector" or "effector moiety" or "effector component" is a molecule that is bound (or linked, or conjugated), either covalently, through a linker or a chemical bond, or noncovalently, through ionic, van der Waals, electrostatic, or hydrogen bonds, to an antibody. The "effector" can be a variety of molecules including, e.g., detection moieties including radioactive compounds, fluorescent compounds, an enzyme or substrate, tags such as epitope tags, a toxin; activatable moieties, a chemotherapeutic agent; a lipase; an antibiotic; or a radioisotope emitting "hard" e.g., beta radiation.

A "labeled nucleic acid probe or oligonucleotide" is one that is bound, either covalently, through a linker or a chemical bond, or noncovalently, through ionic, van der Waals, electrostatic, or hydrogen bonds to a label such that the presence of the probe may be detected by detecting the presence of the label bound to the probe. Alternatively, method using high affinity interactions may achieve the same results where one of a pair of binding partners binds to the other, e.g., biotin, streptavidin.

As used herein a "nucleic acid probe or oligonucleotide" is defined as a nucleic acid capable of binding to a target nucleic acid of complementary sequence through one or more types of chemical bonds, usually through complementary base pairing, usually through hydrogen bond formation. As used herein, a probe may include natural (i.e., A, G, C, or T) or modified bases (7-deazaguanosine, inosine, etc.). In addition, the bases in a probe may be joined by a linkage other than a phosphodiester bond, so long as it does not functionally

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interfere with hybridization. Thus, e.g., probes may be peptide nucleic acids in which the constituent bases are joined by peptide bonds rather than phosphodiester linkages. It will be understood by one of skill in the art that probes may bind target sequences lacking complete complementarity with the probe sequence depending upon the stringency of the hybridization conditions. The probes are preferably directly labeled as with isotopes, chromophores, lumiphores, chromogens, or indirectly labeled such as with biotin to which a streptavidin complex may later bind. By assaying for the presence or absence of the probe, one can detect the presence or absence of the select sequence or subsequence. Diagnosis or prognosis may be based at the genomic level, or at the level of RNA or protein expression.

The term "recombinant" when used with reference, e.g., to a cell, or nucleic acid, protein, or vector, indicates that the cell, nucleic acid, protein or vector, has been modified by the introduction of a heterologous nucleic acid or protein or the alteration of a native nucleic acid or protein, or that the cell is derived from a cell so modified. Thus, e.g., recombinant cells express genes that are not found within the native (non-recombinant) form of the cell or express native genes that are otherwise abnormally expressed, under expressed or not expressed at all. By the term "recombinant nucleic acid" herein is meant nucleic acid, originally formed in vitro, in general, by the manipulation of nucleic acid, e.g., using polymerases and endonucleases, in a form not normally found in nature. In this manner, operably linkage of different sequences is achieved. Thus an isolated nucleic acid, in a linear form, or an expression vector formed in vitro by ligating DNA molecules that are not normally joined, are both considered recombinant for the purposes of this invention. It is understood that once a recombinant nucleic acid is made and reintroduced into a host cell or organism, it will replicate non-recombinantly, i.e., using the in vivo cellular machinery of the host cell rather than in vitro manipulations; however, such nucleic acids, once produced recombinantly, although subsequently replicated non-recombinantly, are still considered recombinant for the purposes of the invention. Similarly, a "recombinant protein" is a protein made using recombinant techniques, i.e., through the expression of a recombinant nucleic acid as depicted above.

The term "heterologous" when used with reference to portions of a nucleic acid indicates that the nucleic acid comprises two or more subsequences that are not normally found in the same relationship to each other in nature. For instance, the nucleic acid is typically recombinantly produced, having two or more sequences, e.g., from unrelated genes

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arranged to make a new functional nucleic acid, e.g., a promoter from one source and a coding region from another source. Similarly, a heterologous protein will often refer to two or more subsequences that are not found in the same relationship to each other in nature (e.g., a fusion protein).

A "promoter" is defined as an array of nucleic acid control sequences that direct transcription of a nucleic acid. As used herein, a promoter includes necessary nucleic acid sequences near the start site of transcription, such as, in the case of a polymerase II type promoter, a TATA element. A promoter also optionally includes distal enhancer or repressor elements, which can be located as much as several thousand base pairs from the start site of transcription. A "constitutive" promoter is a promoter that is active under most environmental and developmental conditions. An "inducible" promoter is a promoter that is active under environmental or developmental regulation. The term "operably linked" refers to a functional linkage between a nucleic acid expression control sequence (such as a promoter, or array of transcription factor binding sites) and a second nucleic acid sequence, wherein the expression control sequence directs transcription of the nucleic acid corresponding to the second sequence.

An "expression vector" is a nucleic acid construct, generated recombinantly or synthetically, with a series of specified nucleic acid elements that permit transcription of a particular nucleic acid in a host cell. The expression vector can be part of a plasmid, virus, or nucleic acid fragment. Typically, the expression vector includes a nucleic acid to be transcribed operably linked to a promoter.

The phrase "selectively (or specifically) hybridizes to" refers to the binding, duplexing, or hybridizing of a molecule only to a particular nucleotide sequence under stringent hybridization conditions when that sequence is present in a complex mixture (e.g., total cellular or library DNA or RNA).

The phrase "stringent hybridization conditions" refers to conditions under which a probe will hybridize to its target subsequence, typically in a complex mixture of nucleic acids, but to no other sequences. Stringent conditions are sequence-dependent and will be different circumstances. Longer sequences hybridize specifically at higher temperatures. An extensive guide to the hybridization of nucleic acids is found "Overview of principles of hybridization and the strategy of nucleic acid assays" in Tijssen (1993) Hybridization with Nucleic Probes (Techniques in Biochemistry and Molecular Biology vol.

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24) Elsevier. Generally, stringent conditions are selected to be about 5-10° C lower than the thermal melting point  $(T_m)$  for the specific sequence at a defined ionic strength pH. The  $T_m$  is the temperature (under defined ionic strength, pH, and nucleic concentration) at which 50% of the probes complementary to the target hybridize to the target sequence at equilibrium (as the target sequences are present in excess, at T<sub>m</sub>, 50% of the probes are occupied at equilibrium). Stringent conditions will be those in which the salt concentration is less than about 1.0 M sodium ion, typically about 0.01 to 1.0 M sodium ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes (e.g., 10 to 50 nucleotides) and at least about 60° C for long probes (e.g., greater than 50 nucleotides). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide. For selective or specific hybridization, a positive signal is at least two times background, preferably 10 times background hybridization. Exemplary stringent hybridization conditions can be as following: 50% formamide, 5x SSC, and 1% SDS, incubating at 42° C, or, 5x SSC, 1% SDS, incubating at 65° C, with wash in 0.2x SSC, and 0.1% SDS at 65° C. For PCR, a temperature of about 36° C is typical for low stringency amplification, although annealing temperatures may vary between about 32° C and 48° C depending on primer length. For high stringency PCR amplification, a temperature of about 62° C is typical, although high stringency annealing temperatures can range from about 50-65° C, depending on the primer length and specificity. Typical cycle conditions for both high and low stringency amplifications include a denaturation phase of 90-95° C for 30-120 sec, an annealing phase lasting 30-120 sec, and an extension phase of about 72° C for 1-2 min. Protocols and guidelines for low and high stringency amplification reactions are provided, e.g., in Innis, et al. (1990) PCR Protocols: A Guide to Methods and Applications Academic Press, N.Y.

Nucleic acids that do not hybridize to each other under stringent conditions are still substantially identical if the polypeptides which they encode are substantially identical. This occurs, e.g., when a copy of a nucleic acid is created using the maximum codon degeneracy permitted by the genetic code. In such cases, the nucleic acids typically hybridize under moderately stringent hybridization conditions. Exemplary "moderately stringent hybridization conditions" include a hybridization in a buffer of 40% formamide, 1 M NaCl, 1% SDS at 37° C, and a wash in 1X SSC at 45° C. A positive hybridization is at least twice

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background. Those of ordinary skill will readily recognize that alternative hybridization and wash conditions can be utilized to provide conditions of similar stringency. Additional guidelines for determining hybridization parameters are provided in numerous references, e.g., Ausubel, et al. (eds. 1991 and supplements) <u>Current Protocols in Molecular Biology</u>

The phrase "functional effects" in the context of assays for testing compounds that modulate activity of a prostate cancer protein includes the determination of a parameter that is indirectly or directly under the influence of the prostate cancer protein or nucleic acid, e.g., a functional, physical, or chemical effect, such as the ability to decrease prostate proliferation (malignant or non-malignant). It includes ligand binding activity; cell growth on soft agar; anchorage dependence; contact inhibition and density limitation of growth; cellular proliferation; cellular transformation; growth factor or serum dependence; tumor specific marker levels; invasiveness into Matrigel; tumor growth and metastasis in vivo; mRNA and protein expression in cells undergoing metastasis, and other characteristics of prostate cancer cells. "Functional effects" include in vitro, in vivo, and ex vivo activities.

By "determining the functional effect" is meant assaying for a compound that increases or decreases a parameter that is indirectly or directly under the influence of a prostate cancer protein sequence, e.g., functional, enzymatic, physical and chemical effects. Such functional effects can be measured by means known to those skilled in the art, e.g., changes in spectroscopic characteristics (e.g., fluorescence, absorbance, refractive index), hydrodynamic (e.g., shape), chromatographic, or solubility properties for the protein, measuring inducible markers or transcriptional activation of the prostate cancer protein; measuring binding activity or binding assays, e.g., binding to antibodies or other ligands, and measuring cellular proliferation. Determination of the functional effect of a compound on prostate cancer can also be performed using prostate cancer assays known to those of skill in the art such as an in vitro assays, e.g., cell growth on soft agar; anchorage dependence; contact inhibition and density limitation of growth; cellular proliferation; cellular transformation; growth factor or serum dependence; tumor specific marker levels; invasiveness into Matrigel; tumor growth and metastasis in vivo; mRNA and protein expression in cells undergoing metastasis, and other characteristics of prostate cancer cells. The functional effects can be evaluated by many means known to those skilled in the art, e.g., microscopy for quantitative or qualitative measures of alterations in morphological features,

measurement of changes in RNA or protein levels for prostate cancer-associated sequences,

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measurement of RNA stability, identification of downstream or reporter gene expression (CAT, luciferase,  $\beta$ -gal, GFP, and the like), e.g., via chemiluminescence, fluorescence, colorimetric reactions, antibody binding, inducible markers, and ligand binding assays.

"Inhibitors", "activators", and "modulators" of prostate cancer polynucleotide and polypeptide sequences are used to refer to activating, inhibitory, or modulating molecules or compounds identified using in vitro and in vivo assays of prostate cancer polynucleotide and polypeptide sequences. Inhibitors are compounds that, e.g., bind to, partially or totally block activity, decrease, prevent, delay activation, inactivate, desensitize, or down regulate the activity or expression of prostate cancer proteins, e.g., antagonists. Antisense nucleic acids may seem to inhibit expression and subsequent function of the protein. "Activators" are compounds that increase, open, activate, facilitate, enhance activation, sensitize, agonize, or up regulate prostate cancer protein activity. Inhibitors, activators, or modulators also include genetically modified versions of prostate cancer proteins, e.g., versions with altered activity, as well as naturally occurring and synthetic ligands, antagonists, agonists, antibodies, small chemical molecules and the like. Such assays for inhibitors and activators include, e.g., expressing the prostate cancer protein in vitro, in cells, or cell membranes, applying putative modulator compounds, and then determining the functional effects on activity, as described above. Activators and inhibitors of prostate cancer can also be identified by incubating prostate cancer cells with the test compound and determining increases or decreases in the expression of 1 or more prostate cancer proteins, e.g., 1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 40, 50 or more prostate cancer proteins, such as prostate cancer proteins encoded by the sequences set out in Tables 1A-4.

Samples or assays comprising prostate cancer proteins that are treated with a potential activator, inhibitor, or modulator are compared to control samples without the inhibitor, activator, or modulator to examine the extent of inhibition. Control samples (untreated with inhibitors) are assigned a relative protein activity value of 100%. Inhibition of a polypeptide is achieved when the activity value relative to the control is about 80%, preferably 50%, more preferably 25-0%. Activation of a prostate cancer polypeptide is achieved when the activity value relative to the control (untreated with activators) is 110%, more preferably 150%, more preferably 200-500% (i.e., two to five fold higher relative to the control), more preferably 1000-3000% higher.

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The phrase "changes in cell growth" refers to a change in cell growth and proliferation characteristics in vitro or in vivo, such as cell viability, formation of foci, anchorage independence, semi-solid or soft agar growth, changes in contact inhibition and density limitation of growth, loss of growth factor or serum requirements, changes in cell morphology, gaining or losing immortalization, gaining or losing tumor specific markers, ability to form or suppress tumors when injected into suitable animal hosts, and/or immortalization of the cell. See, e.g., pp. 231-241 in Freshney (1994) <u>Culture of Animal Cells: A Manual of Basic Technique</u> (3d ed.) Wiley-Liss.

"Tumor cell" refers to precancerous, cancerous, and/or normal cells in a tumor.

"Cancer cells," "transformed" cells, or "transformation" in tissue culture, refers to spontaneous or induced phenotypic changes that do not necessarily involve the uptake of new genetic material. Although transformation can arise from infection with a transforming virus and incorporation of new genomic DNA, or uptake of exogenous DNA, it can also arise spontaneously or following exposure to a carcinogen, thereby mutating an endogenous gene. Transformation is associated with phenotypic changes, such as immortalization of cells, aberrant growth control, nonmorphological changes, and/or malignancy. See, Freshney (2001) Culture of Animal Cells: A Manual of Basic Technique (4th ed.) Wiley-Liss.

"Antibody" refers to a polypeptide comprising a framework region from an immunoglobulin gene or fragments thereof that specifically binds and recognizes an antigen. The recognized immunoglobulin genes include the kappa, lambda, alpha, gamma, delta, epsilon, and mu constant region genes, as well as the myriad immunoglobulin variable region genes. Light chains are classified as either kappa or lambda. Heavy chains are classified as gamma, mu, alpha, delta, or epsilon, which in turn define the immunoglobulin classes, IgG, IgM, IgA, IgD, and IgE, respectively. Typically, the antigen-binding region of an antibody or its functional equivalent will be most critical in specificity and affinity of binding. See Paul (ed. 1999) Fundamental Immunology (4th ed.) Raven.

An exemplary immunoglobulin (antibody) structural unit comprises a tetramer. Each tetramer is composed of two identical pairs of polypeptide chains, each pair having one "light" (about 25 kD) and one "heavy" chain (about 50-70 kD). The N-terminus of each chain defines a variable region of about 100 to 110 or more amino acids primarily responsible for antigen recognition. The terms variable light chain (V<sub>L</sub>) and variable heavy chain (V<sub>H</sub>) refer to these light and heavy chains respectively.

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Antibodies exist, e.g., as intact immunoglobulins or as a number of well-characterized fragments produced by digestion with various peptidases. Thus, e.g., pepsin digests an antibody below the disulfide linkages in the hinge region to produce F(ab)'<sub>2</sub>, a dimer of Fab which itself is a light chain joined to V<sub>H</sub>-C<sub>H</sub>1 by a disulfide bond. The F(ab)'<sub>2</sub> may be reduced under mild conditions to break the disulfide linkage in the hinge region, thereby converting the F(ab)'<sub>2</sub> dimer into an Fab' monomer. The Fab' monomer is essentially Fab with part of the hinge region (see Paul (ed. 1993) <u>Fundamental Immunology</u> (3d ed.) Raven. While various antibody fragments are defined in terms of the digestion of an intact antibody, one of skill will appreciate that such fragments may be synthesized de novo either chemically or by using recombinant DNA methodology. Thus, the term antibody, as used herein, also includes antibody fragments either produced by the modification of whole antibodies, or those synthesized de novo using recombinant DNA methodologies (e.g., single chain Fv) or those identified using phage display libraries (see, e.g., McCafferty, et al.(1990) <u>Nature</u> 348:552-554.

For preparation of antibodies, e.g., recombinant, monoclonal, or polyclonal antibodies, many technique known in the art can be used (see, e.g., Kohler and Milstein (1975) Nature 256:495-497; Kozbor, et al. (1983) Immunology Today 4:72; pp. 77-96 in Cole, et al. (1985) Monoclonal Antibodies and Cancer Therapy Liss; Coligan (1991) Current Protocols in Immunology Lippincott; Harlow and Lane (1988) Antibodies: A Laboratory Manual CSH Press; and Goding (1986) Monoclonal Antibodies: Principles and Practice (2d ed.) Academic Press. Techniques for the production of single chain antibodies (U.S. Patent 4,946,778) can be adapted to produce antibodies to polypeptides of this invention. Also, transgenic mice, or other organisms such as other mammals, may be used to express humanized antibodies. Alternatively, phage display technology can be used to identify antibodies and heteromeric Fab fragments that specifically bind to selected antigens (see, e.g., McCafferty, et al. (1990) Nature 348:552-554; Marks, et al. (1992) Biotechnology 10:779-783).

A "chimeric antibody" is an antibody molecule in which (a) the constant region, or a portion thereof, is altered, replaced or exchanged so that the antigen binding site (variable region) is linked to a constant region of a different or altered class, effector function and/or species, or an entirely different molecule which confers new properties to the chimeric antibody, e.g., an enzyme, toxin, hormone, growth factor, drug, etc.; or (b) the variable

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region, or a portion thereof, is altered, replaced or exchanged with a variable region having a different or altered antigen specificity.

Identification of prostate cancer-associated sequences

In one aspect, the expression levels of genes are determined in different patient samples for which diagnosis information is desired, to provide expression profiles. An expression profile of a particular sample is essentially a "fingerprint" of the state of the sample; while two states may have a particular gene similarly expressed, the evaluation of a number of genes simultaneously allows the generation of a gene expression profile that is characteristic of the state of the cell. That is, normal tissue (e.g., normal prostate or other tissue) may be distinguished from pathological prostate cells, e.g., cancerous or metastatic cancerous tissue of the prostate, or prostate cancer tissue or metastatic prostate cancerous tissue can be compared with tissue samples of prostate and other tissues from surviving cancer patients. By comparing expression profiles of tissue in known different prostate cancer states, information regarding which genes are important (including both up- and down-regulation of genes) in each of these states is obtained.

The identification of sequences that are differentially expressed in prostate cancer versus non-prostate cancer tissue allows the use of this information in a number of ways. For example, a particular treatment regime may be evaluated: does a chemotherapeutic drug act to down-regulate prostate cancer or other proliferative disorders, and thus tumor growth or recurrence, in a particular patient. Alternatively, a treatment step may induce other markers which may be used as targets to destroy tumor cells. Similarly, diagnosis and treatment outcomes may be done or confirmed by comparing patient samples with the known expression profiles. Maliganant disease may be compared to non-malignant conditions. Metastatic tissue can also be analyzed to determine the stage of prostate cancer in the tissue, or origin of primary tumor, e.g., metastasis from a remote primary site. Furthermore, these gene expression profiles (or individual genes) allow screening of drug candidates with an eye to mimicking or altering a particular expression profile; e.g., screening can be done for drugs that suppress the prostate cancer expression profile. This may be done by making biochips comprising sets of the important prostate cancer genes, which can then be used in these screens. These methods can also be done on the protein basis; that is, protein expression levels of the prostate cancer proteins can be evaluated for diagnostic purposes or to screen

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candidate agents. In addition, the prostate cancer nucleic acid sequences can be administered for gene therapy purposes, including the administration of antisense nucleic acids, or the prostate cancer proteins (including antibodies and other modulators thereof) administered as therapeutic drugs.

Thus the present invention provides nucleic acid and protein sequences that are differentially expressed in prostate cancer relative to normal tissues and/or non-malignant disease, or in different types of related diseases, herein termed "prostate cancer sequences." As outlined below, prostate cancer sequences include those that are up-regulated (i.e., expressed at a higher level) in prostate cancer, as well as those that are down-regulated (i.e., expressed at a lower level). In a preferred embodiment, the prostate cancer sequences are from humans; however, as will be appreciated by those in the art, prostate cancer sequences from other organisms may be useful in animal models of disease and drug evaluation; thus, other prostate cancer sequences are provided, from vertebrates, including mammals, including rodents (rats, mice, hamsters, guinea pigs, etc.), primates, farm animals (including sheep, goats, pigs, cows, horses, etc.) and pets, e.g., (dogs, cats, etc.). Prostate cancer sequences from other organisms may be obtained using the techniques outlined below.

Prostate cancer sequences can include both nucleic acid and amino acid sequences. As will be appreciated by those in the art and is more fully outlined below, prostate cancer nucleic acid sequences are useful in a variety of applications, including diagnostic applications, which will detect naturally occurring nucleic acids, as well as screening applications; e.g., biochips comprising nucleic acid probes or PCR microtiter plates with selected probes to the prostate cancer sequences can be generated.

A prostate cancer sequence can be initially identified by substantial nucleic acid and/or amino acid sequence homology to the prostate cancer sequences outlined herein. Such homology can be based upon the overall nucleic acid or amino acid sequence, and is generally determined as outlined below, using either homology programs or hybridization conditions.

For identifying prostate cancer-associated sequences, the prostate cancer screen typically includes comparing genes identified in different tissues, e.g., normal and cancerous tissues, or tumor tissue samples from patients who have metastatic disease vs. non metastatic tissue. Other suitable tissue comparisons include comparing prostate cancer samples with metastatic cancer samples from other cancers, such as lung, breast, gastrointestinal cancers,

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ovarian, etc. Samples of different stages of prostate cancer, e.g., survivor tissue, drug resistant states, and tissue undergoing metastasis, are applied to biochips comprising nucleic acid probes. The samples are first microdissected, if applicable, and treated as is known in the art for the preparation of mRNA. Suitable biochips are commercially available, e.g., from Affymetrix. Gene expression profiles are generated and the data analyzed.

In one embodiment, the genes showing changes in expression as between normal and disease states are compared to genes expressed in other normal tissues, preferably normal prostate, but also including, and not limited to lung, heart, brain, liver, breast, kidney, muscle, colon, small intestine, large intestine, spleen, bone, and placenta. In a preferred embodiment, those genes identified during the prostate cancer screen that are expressed in a significant amount in other tissues are removed from the profile, although in some embodiments, this is not necessary. That is, when screening for drugs, it is usually preferable that the target be disease specific, to minimize possible side effects on other organs were there expression.

In a preferred embodiment, prostate cancer sequences are those that are up-regulated in prostate cancer or related conditions; that is, the expression of these genes is higher in the prostate cancer tissue as compared to non-cancerous tissue. "Up-regulation" as used herein often means at least about a two-fold change, preferably at least about a three fold change, with at least about five-fold or higher being preferred. Another embodiment is directed to sequences up-regulated in non-malignant conditions relative to normal.

Unigene cluster identification numbers and accession numbers herein are for the GenBank sequence database and the sequences of the accession numbers are hereby expressly incorporated by reference. GenBank is known in the art, see, e.g., Benson, et al. (1998) Nucleic Acids Research 26:1-7 and http://www.ncbi.nlm.nih.gov/. Sequences are also available in other databases, e.g., European Molecular Biology Laboratory (EMBL) and DNA Database of Japan (DDBJ). U.S. Patent Application N. 09/687,576 and 09/976,858 (-001-3) further disclose related sequences, compositions, and methods of diagnosis and treatment of prostate cancer and related conditions and are hereby expressly incorporated by reference.

In another preferred embodiment, prostate cancer sequences are those that are downregulated in the prostate cancer; that is, the expression of these genes is lower in prostate cancer tissue as compared to non-cancerous tissue. "Down-regulation" as used herein often

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means at least about a two-fold change, preferably at least about a three fold change, with at least about five-fold or higher being preferred.

### Informatics

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The ability to identify genes that are over or under expressed in prostate cancer can additionally provide high-resolution, high-sensitivity datasets which can be used in the areas of diagnostics, therapeutics, drug development, pharmacogenetics, protein structure, biosensor development, and other related areas. For example, the expression profiles can be used in diagnostic or prognostic evaluation of patients with prostate cancer. Or as another example, subcellular toxicological information can be generated to better direct drug structure and activity correlation (see Anderson, Pharmaceutical Proteomics: Targets, Mechanism, and Function, paper presented at the IBC Proteomics conference, Coronado, CA (June 11-12, 1998)). Subcellular toxicological information can also be utilized in a biological sensor device to predict the likely toxicological effect of chemical exposures and likely tolerable exposure thresholds (see U.S. Patent No. 5,811,231). Similar advantages accrue from datasets relevant to other biomolecules and bioactive agents (e.g., nucleic acids, saccharides, lipids, drugs, and the like).

Thus, in another embodiment, the present invention provides a database that includes at least one set of assay data. The data contained in the database is acquired, e.g., using array analysis either singly or in a library format. The database can be in a form in which data can be maintained and transmitted, but is preferably an electronic database. The electronic database of the invention can be maintained on an electronic device allowing for the storage of and access to the database, such as a personal computer, but is preferably distributed on a wide area network, such as the World Wide Web.

The focus of the present section on databases that include peptide sequence data is for clarity of illustration only. It will be apparent to those of skill in the art that similar databases can be assembled for assay data acquired using an assay of the invention.

The compositions and methods for identifying and/or quantitating the relative and/or absolute abundance of a variety of molecular and macromolecular species from a biological sample undergoing prostate cancer, i.e., the identification of prostate cancer-associated sequences described herein, provide an abundance of information, which can be correlated with pathological conditions, predisposition to disease, drug testing, therapeutic monitoring,

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gene-disease causal linkages, identification of correlates of immunity and physiological status, among others. Although the data generated from the assays of the invention is suited for manual review and analysis, in a preferred embodiment, prior data processing using high-speed computers is utilized.

An array of methods for indexing and retrieving biomolecular information is known in the art. For example, U.S. Patents 6,023,659 and 5,966,712 disclose a relational database system for storing biomolecular sequence information in a manner that allows sequences to be catalogued and searched according to one or more protein function hierarchies. U.S. Patent 5,953,727 discloses a relational database having sequence records containing information in a format that allows a collection of partial-length DNA sequences to be catalogued and searched according to association with one or more sequencing projects for obtaining full-length sequences from the collection of partial length sequences. U.S. Patent 5,706,498 discloses a gene database retrieval system for making a retrieval of a gene sequence similar to a sequence data item in a gene database based on the degree of similarity between a key sequence and a target sequence. U.S. Patent 5,538,897 discloses a method using mass spectroscopy fragmentation patterns of peptides to identify amino acid sequences in computer databases by comparison of predicted mass spectra with experimentally-derived mass spectra using a closeness-of-fit measure. U.S. Patent 5,926,818 discloses a multidimensional database comprising a functionality for multi-dimensional data analysis described as on-line analytical processing (OLAP), which entails the consolidation of projected and actual data according to more than one consolidation path or dimension. U.S. Patent 5,295,261 reports a hybrid database structure in which the fields of each database record are divided into two classes, navigational and informational data, with navigational fields stored in a hierarchical topological map which can be viewed as a tree structure or as the merger of two or more such tree structures.

See also Mount, et al. (2001) <u>Bioinformatics</u> CSH Press; Durbin, et al. (eds. 1999) <u>Biological Sequence Analysis: Probabilistic Models of Proteins and Nucleic Acids</u> Cambridge Univ. Press; Baxevanis and Oeullette (eds., 1998) <u>Bioinformatics: A Practical Guide to the Analysis of Genes and Proteins</u> Wiley-Liss; Rashidi and Buehler (1999) <u>Bioinformatics: Basic Applications in Biological Science and Medicine</u> CRC Press; Setubal, et al. (eds. 1997) <u>Introduction to Computational Molecular Biology</u> Brooks/Cole; Misener and Krawetz (eds. 2000) <u>Bioinformatics: Methods and Protocols Human Press; Higgins and</u>

Taylor (eds. 2000) <u>Bioinformatics: Sequence, Structure, and Databanks: A Practical Approach</u> Oxford Univ. Press; Brown (2001) <u>Bioinformatics: A Biologist's Guide to Biocomputing and the Internet Eaton Pub; Han and Kamber (2000) <u>Data Mining: Concepts and Techniques</u> Kaufmann Pub.; and Waterman (1995) <u>Introduction to Computational Biology: Maps, Sequences, and Genomes</u> Chap and Hall.</u>

The present invention provides a computer database comprising a computer and software for storing in computer-retrievable form assay data records cross-tabulated, e.g., with data specifying the source of the target-containing sample from which each sequence specificity record was obtained.

In an exemplary embodiment, at least one of the sources of target-containing sample is from a control tissue sample known to be free of pathological disorders. In a variation, at least one of the sources is a known pathological tissue specimen, e.g., a neoplastic lesion or another tissue specimen to be analyzed for prostate cancer. In another variation, the assay records cross-tabulate one or more of the following parameters for each target species in a sample: (1) a unique identification code, which can include, e.g., a target molecular structure and/or characteristic separation coordinate (e.g., electrophoretic coordinates); (2) sample source; and (3) absolute and/or relative quantity of the target species present in the sample.

The invention also provides for the storage and retrieval of a collection of target data in a computer data storage apparatus, which can include magnetic disks, optical disks, magneto-optical disks, DRAM, SRAM, SGRAM, SDRAM, RDRAM, DDR RAM, magnetic bubble memory devices, and other data storage devices, including CPU registers and on-CPU data storage arrays. Typically, the target data records are stored as a bit pattern in an array of magnetic domains on a magnetizable medium or as an array of charge states or transistor gate states, such as an array of cells in a DRAM device (e.g., each cell comprised of a transistor and a charge storage area, which may be on the transistor). In one embodiment, the invention provides such storage devices, and computer systems built therewith, comprising a bit pattern encoding a protein expression fingerprint record comprising unique identifiers for at least 10 target data records cross-tabulated with target source.

When the target is a peptide or nucleic acid, the invention preferably provides a method for identifying related peptide or nucleic acid sequences, comprising performing a computerized comparison between a peptide or nucleic acid sequence assay record stored in or retrieved from a computer storage device or database and at least one other sequence. The

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comparison can include a sequence analysis or comparison algorithm or computer program embodiment thereof (e.g., FASTA, TFASTA, GAP, BESTFIT) and/or the comparison may be of the relative amount of a peptide or nucleic acid sequence in a pool of sequences determined from a polypeptide or nucleic acid sample of a specimen.

The invention also preferably provides a magnetic disk, such as an IBM-compatible (DOS, Windows, Windows95/98/2000, Windows NT, OS/2) or other format (e.g., Linux, SunOS, Solaris, AIX, SCO Unix, VMS, MV, Macintosh, etc.) floppy diskette or hard (fixed, Winchester) disk drive, comprising a bit pattern encoding data from an assay of the invention in a file format suitable for retrieval and processing in a computerized sequence analysis, comparison, or relative quantitation method.

The invention also provides a network, comprising a plurality of computing devices linked via a data link, such as an Ethernet cable (coax or 10BaseT), telephone line, ISDN line, wireless network, optical fiber, or other suitable signal transmission medium, whereby at least one network device (e.g., computer, disk array, etc.) comprises a pattern of magnetic domains (e.g., magnetic disk) and/or charge domains (e.g., an array of DRAM cells) composing a bit pattern encoding data acquired from an assay of the invention.

The invention also provides a method for transmitting assay data that includes generating an electronic signal on an electronic communications device, such as a modern, ISDN terminal adapter, DSL, cable modern, ATM switch, or the like, wherein the signal includes (in native or encrypted format) a bit pattern encoding data from an assay or a database comprising a plurality of assay results obtained by the method of the invention.

In a preferred embodiment, the invention provides a computer system for comparing a query target to a database containing an array of data structures, such as an assay result obtained by the method of the invention, and ranking database targets based on the degree of identity and gap weight to the target data. A central processor is preferably initialized to load and execute the computer program for alignment and/or comparison of the assay results. Data for a query target is entered into the central processor via an I/O device. Execution of the computer program results in the central processor retrieving the assay data from the data file, which comprises a binary description of an assay result.

The target data or record and the computer program can be transferred to secondary memory, which is typically random access memory (e.g., DRAM, SRAM, SGRAM, or SDRAM). Targets are ranked according to the degree of correspondence between a selected

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assay characteristic (e.g., binding to a selected affinity moiety) and the same characteristic of the query target and results are output via an I/O device. For example, a central processor can be a conventional computer (e.g., Intel Pentium, PowerPC, Alpha, PA-8000, SPARC, MIPS 4400, MIPS 10000, VAX, etc.); a program can be a commercial or public domain molecular biology software package (e.g., UWGCG Sequence Analysis Software, Darwin); a data file can be an optical or magnetic disk, a data server, a memory device (e.g., DRAM, SRAM, SGRAM, SDRAM, EPROM, bubble memory, flash memory, etc.); an I/O device can be a terminal comprising a video display and a keyboard, a modem, an ISDN terminal adapter, an Ethernet port, a punched card reader, a magnetic strip reader, or other suitable I/O device.

The invention also preferably provides the use of a computer system, such as that described above, which comprises: (1) a computer; (2) a stored bit pattern encoding a collection of peptide sequence specificity records obtained by the methods of the invention, which may be stored in the computer; (3) a comparison target, such as a query target; and (4) a program for alignment and comparison, typically with rank-ordering of comparison results on the basis of computed similarity values.

# Characteristics of prostate cancer-associated proteins

Prostate cancer proteins of the present invention may be classified as secreted proteins, transmembrane proteins, or intracellular proteins. In one embodiment, the prostate cancer protein is an intracellular protein. Intracellular proteins may be found in the cytoplasm and/or in the nucleus. Intracellular proteins are involved in all aspects of cellular function and replication (including, e.g., signaling pathways); aberrant expression of such proteins often results in unregulated or disregulated cellular processes (see, e.g., Alberts (ed. 1994) Molecular Biology of the Cell (3d ed.) Garland. For example, many intracellular proteins have enzymatic activity such as protein kinase activity, protein phosphatase activity, protease activity, nucleotide cyclase activity, polymerase activity and the like. Intracellular proteins also serve as docking proteins that are involved in organizing complexes of proteins, or targeting proteins to various subcellular localizations, and are involved in maintaining the structural integrity of organelles.

An increasingly appreciated concept in characterizing proteins is the presence in the proteins of one or more structural motifs for which defined functions have been attributed. In

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addition to the highly conserved sequences found in the enzymatic domain of proteins, highly conserved sequences have been identified in proteins that are involved in protein-protein interaction. For example, Src-homology-2 (SH2) domains bind tyrosine-phosphorylated targets in a sequence dependent manner. PTB domains, which are distinct from SH2 domains, also bind tyrosine phosphorylated targets. SH3 domains bind to proline-rich targets. In addition, PH domains, tetratricopeptide repeats and WD domains to name only a few, have been shown to mediate protein-protein interactions. Some of these may also be involved in binding to phospholipids or other second messengers. As will be appreciated by one of ordinary skill in the art, these motifs can be identified on the basis of amino acid sequence; thus, an analysis of the sequence of proteins may provide insight into both the enzymatic potential of the molecule and/or molecules with which the protein may associate. One useful database is Pfam (protein families), which is a large collection of multiple sequence alignments and hidden Markov models covering many common protein domains. Versions are available via the internet from Washington University in St. Louis, the Sanger Center in England, and the Karolinska Institute in Sweden (see, e.g., Bateman, et al. (2000) Nuc. Acids Res. 28:263-266; Sonnhammer, et al. (1997) Proteins 28:405-420; Bateman, et al. (1999) Nuc. Acids Res. 27:260-262; and Sonnhammer, et al. (1998) Nuc. Acids Res. 26:320-322.

In another embodiment, the prostate cancer sequences are transmembrane proteins. Transmembrane proteins are molecules that span a phospholipid bilayer of a cell. They may have an intracellular domain, an extracellular domain, or both. The intracellular domains of such proteins may have a number of functions including those already described for intracellular proteins. For example, the intracellular domain may have enzymatic activity and/or may serve as a binding site for additional proteins. Frequently the intracellular domain of transmembrane proteins serves both roles. For example certain receptor tyrosine kinases have both protein kinase activity and SH2 domains. In addition, autophosphorylation of tyrosines on the receptor molecule itself, creates binding sites for additional SH2 domain containing proteins.

Transmembrane proteins may contain from one to many transmembrane domains.

For example, receptor tyrosine kinases, certain cytokine receptors, receptor guanylyl cyclases and receptor serine/threonine protein kinases contain a single transmembrane domain.

However, various other proteins including channels and adenylyl cyclases contain numerous

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transmembrane domains. Many important cell surface receptors such as G protein coupled receptors (GPCRs) are classified as "seven transmembrane domain" proteins, as they contain 7 membrane spanning regions. Characteristics of transmembrane domains include approximately 17 consecutive hydrophobic amino acids that may be followed by charged amino acids. Therefore, upon analysis of the amino acid sequence of a particular protein, the localization and number of transmembrane domains within the protein may be predicted (see, e.g., PSORT web site http://psort.nibb.ac.jp/). Important transmembrane protein receptors include, but are not limited to the insulin receptor, insulin-like growth factor receptor, human growth hormone receptor, glucose transporters, transferrin receptor, epidermal growth factor receptor, low density lipoprotein receptor, epidermal growth factor receptor, leptin receptor, and interleukin receptors, e.g., IL-1 receptor, IL-2 receptor, etc.

The extracellular domains of transmembrane proteins are diverse; however, conserved motifs are found repeatedly among various extracellular domains. Conserved structure and/or functions have been ascribed to different extracellular motifs. Many extracellular domains are involved in binding to other molecules. In one aspect, extracellular domains are found on receptors. Factors that bind the receptor domain include circulating ligands, which may be peptides, proteins, or small molecules such as adenosine and the like. For example, growth factors such as EGF, FGF, and PDGF are circulating growth factors that bind to their cognate receptors to initiate a variety of cellular responses. Other factors include cytokines, mitogenic factors, neurotrophic factors and the like. Extracellular domains also bind to cell-associated molecules. In this respect, they mediate cell-cell interactions. Cell-associated ligands can be tethered to the cell, e.g., via a glycosylphosphatidylinositol (GPI) anchor, or may themselves be transmembrane proteins. Extracellular domains also associate with the extracellular matrix and contribute to the maintenance of the cell structure.

Prostate cancer proteins that are transmembrane are particularly preferred in the present invention as they are readily accessible targets for immunotherapeutics, as are described herein. In addition, as outlined below, transmembrane proteins can be also useful in imaging modalities. Antibodies may be used to label such readily accessible proteins in situ. Alternatively, antibodies can also label intracellular proteins, in which case samples are typically permeablized to provide access to intracellular proteins.. In addition, some membrane proteins can be processed to release a soluble protein, or to expose a residual

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fragment. Released soluble proteins may be useful diagnostic markers, processed residual protein fragments may be useful prostate markers of disease.

It will also be appreciated by those in the art that a transmembrane protein can be made soluble by removing transmembrane sequences, e.g., through recombinant methods. Furthermore, transmembrane proteins that have been made soluble can be made to be secreted through recombinant means by adding an appropriate signal sequence.

In another embodiment, the prostate cancer proteins are secreted proteins; the secretion of which can be either constitutive or regulated. These proteins may have a signal peptide or signal sequence that targets the molecule to the secretory pathway. Secreted proteins are involved in numerous physiological events; by virtue of their circulating nature, they often serve to transmit signals to various other cell types. The secreted protein may function in an autocrine manner (acting on the cell that secreted the factor), a paracrine manner (acting on cells in close proximity to the cell that secreted the factor), an endocrine manner (acting on cells at a distance, e.g., secretion into the blood stream), or an exocrine manner (secretion, e.g., through a duct or to adjacent epithelial surface as sweat glands, sebaceous glands, pancreatic ducts, lacrimal glands, mammary glands, sax producing glands of the ear, etc.). Thus secreted molecules find use in modulating or altering numerous aspects of physiology. Prostate cancer proteins that are secreted proteins are particularly preferred in the present invention as they serve as good targets for diagnostic markers, e.g., for blood, plasma, serum, or stool tests. Those which are enzymes may be antibody or small molecule targets. Others may be useful as vaccine targets, e.g., via CTL mechanisms.

Use of prostate cancer nucleic acids

As described above, prostate cancer sequence is initially identified by substantial nucleic acid and/or amino acid sequence homology or linkage to the prostate cancer sequences outlined herein. Such homology can be based upon the overall nucleic acid or amino acid sequence, and is generally determined as outlined below, using either homology programs or hybridization conditions. Typically, linked sequences on a mRNA are found on the same molecule.

The prostate cancer nucleic acid sequences of the invention, e.g., the sequences in Tables 1A-4, can be fragments of larger genes, i.e., they are nucleic acid segments. "Genes" in this context includes coding regions, non-coding regions, and mixtures of coding and non-

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coding regions. Accordingly, as will be appreciated by those in the art, using the sequences provided herein, extended sequences, in either direction, of the prostate cancer genes can be obtained, using techniques well known in the art for cloning either longer sequences or the full length sequences; see Ausubel, et al., supra. Much can be done by informatics and many sequences can be clustered to include multiple sequences corresponding to a single gene, e.g., systems such as UniGene (see, http://www.ncbi.nlm.nih.gov/UniGene/).

Once the prostate cancer nucleic acid is identified, it can be cloned and, if necessary, its constituent parts recombined to form the entire prostate cancer nucleic acid coding regions or the entire mRNA sequence. Once isolated from its natural source, e.g., contained within a plasmid or other vector or excised therefrom as a linear nucleic acid segment, the recombinant prostate cancer nucleic acid can be further-used as a probe to identify and isolate other prostate cancer nucleic acids, e.g., extended coding regions. It can also be used as a "precursor" nucleic acid to make modified or variant prostate cancer nucleic acids and proteins.

The prostate cancer nucleic acids of the present invention are used in several ways. In a first embodiment, nucleic acid probes to the prostate cancer nucleic acids are made and attached to biochips to be used in screening and diagnostic methods, as outlined below, or for administration, e.g., for gene therapy, vaccine, and/or antisense applications. Alternatively, the prostate cancer nucleic acids that include coding regions of prostate cancer proteins can be put into expression vectors for the expression of prostate cancer proteins, again for screening purposes or for administration to a patient.

In a preferred embodiment, nucleic acid probes to prostate cancer nucleic acids (both the nucleic acid sequences outlined in the figures and/or the complements thereof) are made. The nucleic acid probes attached to the biochip are designed to be substantially complementary to the prostate cancer nucleic acids, i.e., the target sequence (either the target sequence of the sample or to other probe sequences, e.g., in sandwich assays), such that hybridization of the target sequence and the probes of the present invention occurs. As outlined below, this complementarity need not be perfect; there may be base pair mismatches which will interfere with hybridization between the target sequence and the single stranded nucleic acids of the present invention. However, if the number of mutations is so great that no hybridization can occur under even the least stringent of hybridization conditions, the sequence is not a complementary target sequence. Thus, by "substantially complementary"

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herein is meant that the probes are sufficiently complementary to the target sequences to hybridize under normal reaction conditions, particularly high stringency conditions, as outlined herein.

A nucleic acid probe is generally single stranded but can be partially single and partially double stranded. The strandedness of the probe is dictated by the structure, composition, and properties of the target sequence. In general, the nucleic acid probes range from about 8 to about 100 bases long, with from about 10 to about 80 bases being preferred, and from about 30 to about 50 bases being particularly preferred. That is, generally whole genes are not used. In some embodiments, much longer nucleic acids can be used, up to hundreds of bases.

In a preferred embodiment, more than one probe per sequence is used, with either overlapping probes or probes to different sections of the target being used. That is, two, three, four or more probes, with three being preferred, are used to build in a redundancy for a particular target. The probes can be overlapping (i.e., have some sequence in common), or separate. In some cases, PCR primers may be used to amplify signal for higher sensitivity.

As will be appreciated by those in the art, nucleic acids can be attached or immobilized to a solid support in a wide variety of ways. By "immobilized" and grammatical equivalents herein is meant the association or binding between the nucleic acid probe and the solid support is sufficient to be stable under the conditions of binding, washing, analysis, and removal as outlined below. The binding can typically be covalent or non-covalent. By "non-covalent binding" and grammatical equivalents herein is meant one or more of electrostatic, hydrophilic, and hydrophobic interactions. Included in non-covalent binding is the covalent attachment of a molecule, such as, streptavidin to the support and the non-covalent binding of the biotinylated probe to the streptavidin. By "covalent binding" and grammatical equivalents herein is meant that the two moieties, the solid support and the probe, are attached by at least one bond, including sigma bonds, pi bonds and coordination bonds. Covalent bonds can be formed directly between the probe and the solid support or can be formed by a cross linker or by inclusion of a specific reactive group on either the solid support or the probe or both molecules. Immobilization may also involve a combination of covalent and non-covalent interactions.

In general, the probes are attached to the biochip in a wide variety of ways, as will be appreciated by those in the art. As described herein, the nucleic acids can either be

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synthesized first, with subsequent attachment to the biochip, or can be directly synthesized on the biochip.

The biochip comprises a suitable solid substrate. By "substrate" or "solid support" or other grammatical equivalents herein is meant a material that can be modified to contain discrete individual sites appropriate for the attachment or association of the nucleic acid probes and is amenable to at least one detection method. As will be appreciated by those in the art, the number of possible substrates are very large, and include, but are not limited to, glass and modified or functionalized glass, plastics (including acrylics, polystyrene and copolymers of styrene and other materials, polypropylene, polyethylene, polybutylene, polyurethanes, TeflonJ, etc.), polysaccharides, nylon or nitrocellulose, resins, silica or silicabased materials including silicon and modified silicon, carbon, metals, inorganic glasses, plastics, etc. In general, the substrates allow optical detection and do not appreciably fluoresce. A preferred substrate is described in WO0055627, herein incorporated by reference in its entirety.

Generally the substrate is planar, although as will be appreciated by those in the art, other configurations of substrates may be used as well. For example, the probes may be placed on the inside surface of a tube, for flow-through sample analysis to minimize sample volume. Similarly, the substrate may be flexible, such as a flexible foam, including closed cell foams made of particular plastics.

In a preferred embodiment, the surface of the biochip and the probe may be derivatized with chemical functional groups for subsequent attachment of the two. Thus, e.g., the biochip is derivatized with a chemical functional group including, but not limited to, amino groups, carboxy groups, oxo groups and thiol groups, with amino groups being particularly preferred. Using these functional groups, the probes can be attached using functional groups on the probes. For example, nucleic acids containing amino groups can be attached to surfaces comprising amino groups, e.g., using linkers as are known in the art; e.g., homo-or hetero-bifunctional linkers as are well known (see 1994 Pierce Chemical Company catalog, technical section on cross-linkers, pages 155-200). In addition, in some cases, additional linkers, such as alkyl groups (including substituted and heteroalkyl groups) may be used.

In this embodiment, oligonucleotides are synthesized as is known in the art, and then attached to the surface of the solid support. As will be appreciated by those skilled in the art,

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either the 5' or 3' terminus may be attached to the solid support, or attachment may be via an internal nucleoside.

In another embodiment, the immobilization to the solid support may be very strong, yet non-covalent. For example, biotinylated oligonucleotides can be made, which bind to surfaces covalently coated with streptavidin, resulting in attachment.

Alternatively, the oligonucleotides may be synthesized on the surface, as is known in the art. For example, photoactivation techniques utilizing photopolymerization compounds and techniques are used. In a preferred embodiment, the nucleic acids can be synthesized in situ, using well known photolithographic techniques, such as those described in WO 95/25116; WO 95/35505; U.S. Patent Nos. 5,700,637 and 5,445,934; and references cited within, all of which are expressly incorporated by reference; these methods of attachment form the basis of the Affymetrix GeneChip<sup>TM</sup> technology.

Often, amplification-based assays are performed to measure the expression level of prostate cancer-associated sequences. These assays are typically performed in conjunction with reverse transcription. In such assays, a prostate cancer-associated nucleic acid sequence acts as a template in an amplification reaction (e.g., Polymerase Chain Reaction, or PCR). In a quantitative amplification, the amount of amplification product will be proportional to the amount of template in the original sample. Comparison to appropriate controls provides a measure of the amount of prostate cancer-associated RNA. Methods of quantitative amplification are well known to those of skill in the art. Detailed protocols for quantitative PCR are provided, e.g., in Innis, et al. (1990) PCR Protocols: A Guide to Methods and Applications Academic Press.

In some embodiments, a TaqMan based assay is used to measure expression. TaqMan based assays use a fluorogenic oligonucleotide probe that contains a 5' fluorescent dye and a 3' quenching agent. The probe hybridizes to a PCR product, but cannot itself be extended due to a blocking agent at the 3' end. When the PCR product is amplified in subsequent cycles, the 5' nuclease activity of the polymerase, e.g., AmpliTaq, results in the cleavage of the TaqMan probe. This cleavage separates the 5' fluorescent dye and the 3' quenching agent, thereby resulting in an increase in fluorescence as a function of amplification (see, e.g., literature provided by Perkin-Elmer, e.g., www2.perkin-elmer.com).

Other suitable amplification methods include, but are not limited to, ligase chain reaction (LCR) (see Wu and Wallace (1989) Genomics 4:560-569, Landegren, et al. (1988)

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Science 241:1077-1080, and Barringer, et al. (1990) Gene 89:117-122), transcription amplification (Kwoh, et al. (1989) Proc. Natl. Acad. Sci. USA 86:1173-1177), self-sustained sequence replication (Guatelli, et al. (1990) Proc. Nat. Acad. Sci. USA 87:1874-1878), dot PCR, and linker adapter PCR, etc.

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Expression of prostate cancer proteins from nucleic acids

In a preferred embodiment, prostate cancer nucleic acids, e.g., encoding prostate cancer proteins are used to make a variety of expression vectors to express prostate cancer proteins which can then be used in screening assays, as described below. Expression vectors and recombinant DNA technology are well known to those of skill in the art (see, e.g., Ausubel, supra, and Fernandez and Hoeffler (eds. 1999) Gene Expression Systems Academic Press) and are used to express proteins. The expression vectors may be either self-replicating extrachromosomal vectors or vectors which integrate into a host genome. Generally, these expression vectors include transcriptional and translational regulatory nucleic acid operably linked to the nucleic acid encoding the prostate cancer protein. The term "control sequences" refers to DNA sequences used for the expression of an operably linked coding sequence in a particular host organism. Control sequences that are suitable for prokaryotes, e.g., include a promoter, optionally an operator sequence, and a ribosome binding site. Eukaryotic cells are known to utilize promoters, polyadenylation signals, and enhancers.

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Nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the sequence; a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation, and sequences may be operably linked when they are physically linked on the same molecule. Generally, "operably linked" means that the DNA sequences being linked are contiguous, and, in the case of a secretory leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is typically accomplished by ligation at convenient restriction sites. If such sites do not exist, synthetic oligonucleotide adaptors or linkers are used in accordance with conventional practice. Transcriptional and translational regulatory nucleic acid will generally be appropriate to the host cell used to express the prostate cancer protein.

Numerous types of appropriate expression vectors, and suitable regulatory sequences are known in the art for a variety of host cells.

In general, transcriptional and translational regulatory sequences may include, but are not limited to, promoter sequences, ribosomal binding sites, transcriptional start and stop sequences, translational start and stop sequences, and enhancer or activator sequences. In a preferred embodiment, the regulatory sequences include a promoter and transcriptional start and stop sequences.

Promoter sequences encode either constitutive or inducible promoters. The promoters may be either naturally occurring promoters or hybrid promoters. Hybrid promoters, which combine elements of more than one promoter, are also known in the art, and are useful in the present invention.

In addition, an expression vector may comprise additional elements. For example, the expression vector may have two replication systems, thus allowing it to be maintained in two organisms, e.g., in mammalian or insect cells for expression and in a prokaryotic host for cloning and amplification. Furthermore, for integrating expression vectors, the expression vector contains at least one sequence homologous to the host cell genome, and preferably two homologous sequences which flank the expression construct. The integrating vector may be directed to a specific locus in the host cell by selecting the appropriate homologous sequence for inclusion in the vector. Constructs for integrating vectors are well known in the art (e.g., Fernandez and Hoeffler, supra).

In addition, in a preferred embodiment, the expression vector contains a selectable marker gene to allow the selection of transformed host cells. Selection genes are well known in the art and will vary with the host cell used.

The prostate cancer proteins of the present invention are produced by culturing a host cell transformed with an expression vector containing nucleic acid encoding a prostate cancer protein, under the appropriate conditions to induce or cause expression of the prostate cancer protein. Conditions appropriate for prostate cancer protein expression will vary with the choice of the expression vector and the host cell, and will be easily ascertained by one skilled in the art through routine experimentation or optimization. For example, the use of constitutive promoters in the expression vector will require optimizing the growth and proliferation of the host cell, while the use of an inducible promoter requires the appropriate growth conditions for induction. In addition, in some embodiments, the timing of the harvest

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is important. For example, the baculoviral systems used in insect cell expression are lytic viruses, and thus harvest time selection can be crucial for product yield.

Appropriate host cells include yeast, bacteria, archaebacteria, fungi, and insect and animal cells, including mammalian cells. Of particular interest are Saccharomyces cerevisiae and other yeasts, E. coli, Bacillus subtilis, Sf9 cells, C129 cells, 293 cells, Neurospora, BHK, CHO, COS, HeLa cells, HUVEC (human umbilical vein endothelial cells), THP1 cells (a macrophage cell line) and various other human cells and cell lines.

In a preferred embodiment, the prostate cancer proteins are expressed in mammalian cells. Mammalian expression systems are also known in the art, and include retroviral and adenoviral systems. One expression vector system is a retroviral vector system such as is generally described in PCT/US97/01019 and PCT/US97/01048, both of which are hereby expressly incorporated by reference. Of particular use as mammalian promoters are the promoters from mammalian viral genes, since the viral genes are often highly expressed and have a broad host range. Examples include the SV40 early promoter, mouse mammary tumor virus LTR promoter, adenovirus major late promoter, herpes simplex virus promoter, and the CMV promoter (see, e.g., Fernandez and Hoeffler, supra). Typically, transcription termination and polyadenylation sequences recognized by mammalian cells are regulatory regions located 3' to the translation stop codon and thus, together with the promoter elements, flank the coding sequence. Examples of transcription terminator and polyadenylation signals include those derived form SV40.

The methods of introducing exogenous nucleic acid into mammalian hosts, as well as other hosts, is well known in the art, and will vary with the host cell used. Techniques include dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, viral infection, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei.

In a preferred embodiment, prostate cancer proteins are expressed in bacterial systems. Bacterial expression systems are well known in the art. Promoters from bacteriophage may also be used and are known in the art. In addition, synthetic promoters and hybrid promoters are also useful; e.g., the tac promoter is a hybrid of the trp and lac promoter sequences. Furthermore, a bacterial promoter can include naturally occurring promoters of non-bacterial origin that have the ability to bind bacterial RNA polymerase and initiate transcription. In addition to a functioning promoter sequence, an efficient ribosome

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binding site is desirable. The expression vector may also include a signal peptide sequence that provides for secretion of the prostate cancer protein in bacteria. The protein is either secreted into the growth media (gram-positive bacteria) or into the periplasmic space, located between the inner and outer membrane of the cell (gram-negative bacteria). The bacterial expression vector may also include a selectable marker gene to allow for the selection of bacterial strains that have been transformed. Suitable selection genes include genes which render the bacteria resistant to drugs such as ampicillin, chloramphenicol, erythromycin, kanamycin, neomycin and tetracycline. Selectable markers also include biosynthetic genes, such as those in the histidine, tryptophan and leucine biosynthetic pathways. These components are assembled into expression vectors. Expression vectors for bacteria are well known in the art, and include vectors for Bacillus subtilis, E. coli, Streptococcus cremoris, and Streptococcus lividans, among others (e.g., Fernandez and Hoeffler, supra). The bacterial expression vectors are transformed into bacterial host cells using techniques well known in the art, such as calcium chloride treatment, electroporation, and others.

In one embodiment, prostate cancer proteins are produced in insect cells. Expression vectors for the transformation of insect cells, and in particular, baculovirus-based expression vectors, are well known in the art.

In a preferred embodiment, prostate cancer protein is produced in yeast cells. Yeast expression systems are well known in the art, and include expression vectors for Saccharomyces cerevisiae, Candida albicans and C. maltosa, Hansenula polymorpha, Kluyveromyces fragilis and K. lactis, Pichia guillerimondii and P. pastoris, Schizosaccharomyces pombe, and Yarrowia lipolytica.

The prostate cancer protein may also be made as a fusion protein, using techniques well known in the art. Thus, e.g., for the creation of monoclonal antibodies, if the desired epitope is small, the prostate cancer protein may be fused to a carrier protein to form an immunogen. Alternatively, the prostate cancer protein may be made as a fusion protein to increase expression, or for other reasons. For example, when the prostate cancer protein is a prostate cancer peptide, the nucleic acid encoding the peptide may be linked to other nucleic acid for expression purposes.

In a preferred embodiment, the prostate cancer protein is purified or isolated after expression. Prostate cancer proteins may be isolated or purified in a variety of ways known to those skilled in the art depending on what other components are present in the sample.

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Standard purification methods include electrophoretic, molecular, immunological and chromatographic techniques, including ion exchange, hydrophobic, affinity, and reverse-phase HPLC chromatography, and chromatofocusing. For example, the prostate cancer protein may be purified using a standard anti-prostate cancer protein antibody column. Ultrafiltration and diafiltration techniques, in conjunction with protein concentration, are also useful. For general guidance in suitable purification techniques, see Scopes (1982) Protein Purification Springer-Verlag. The degree of purification necessary will vary depending on the use of the prostate cancer protein. In some instances no purification will be necessary.

Once expressed and purified if necessary, the prostate cancer proteins and nucleic acids are useful in a number of applications. They may be used as immunoselection reagents, as vaccine reagents, as screening agents, etc.

## Variants of prostate cancer proteins

In one embodiment, the prostate cancer proteins are derivative or variant prostate cancer proteins as compared to the wild-type sequence. That is, as outlined more fully below, the derivative prostate cancer peptide will often contain at least one amino acid substitution, deletion or insertion, with amino acid substitutions being particularly preferred. The amino acid substitution, insertion, or deletion may occur at most any residue within the prostate cancer peptide.

Also included within one embodiment of prostate cancer proteins of the present invention are amino acid sequence variants. These variants typically fall into one or more of three classes: substitutional, insertional, or deletional variants. These variants ordinarily are prepared by site specific mutagenesis of nucleotides in the DNA encoding the prostate cancer protein, using cassette or PCR mutagenesis or other techniques well known in the art, to produce DNA encoding the variant, and thereafter expressing the DNA in recombinant cell culture as outlined above. However, variant prostate cancer protein fragments having up to about 100-150 residues may be prepared by in vitro synthesis using established techniques. Amino acid sequence variants are characterized by the predetermined nature of the variation, a feature that sets them apart from naturally occurring allelic or interspecies variation of the prostate cancer protein amino acid sequence. The variants typically exhibit the same qualitative biological activity as the naturally occurring analogue, although variants can also be selected which have modified characteristics as will be more fully outlined below.

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While the site or region for introducing an amino acid sequence variation is predetermined, the mutation per se need not be predetermined. For example, in order to optimize the performance of a mutation at a given site, random mutagenesis may be conducted at the target codon or region and the expressed prostate cancer variants screened for the optimal combination of desired activity. Techniques for making substitution mutations at predetermined sites in DNA having a known sequence are well known, e.g., M13 primer mutagenesis and PCR mutagenesis. Screening of the mutants is done using assays of prostate cancer protein activities.

Amino acid substitutions are typically of single residues; insertions usually will be on the order of from about 1 to 20 amino acids, although considerably larger insertions may be tolerated. Deletions range from about 1 to about 20 residues, although in some cases deletions may be much larger.

Substitutions, deletions, insertions or a combination thereof may be used to arrive at a final derivative. Generally these changes are done on a few amino acids to minimize the alteration of the molecule. However, larger changes may be tolerated in certain circumstances. When small alterations in the characteristics of the prostate cancer protein are desired, substitutions are generally made in accordance with the amino acid substitution relationships provided in the definition section.

The variants typically exhibit the same qualitative biological activity and will elicit the same immune response as the naturally-occurring analog, although variants also are selected to modify the characteristics of the prostate cancer proteins as needed. Alternatively, the variant may be designed such that the biological activity of the prostate cancer protein is altered. For example, glycosylation sites may be altered or removed.

Substitutions that are less conservative than those described above. For example, substitutions may be made which more significantly affect: the structure of the polypeptide backbone in the area of the alteration, for example the alpha-helical or beta-sheet structure; the charge or hydrophobicity of the molecule at the target site; or the bulk of the side chain. The substitutions which in general are expected to produce the greatest changes in the polypeptide's properties are those in which (a) a hydrophilic residue, e.g., serinyl or threonyl is substituted for (or by) a hydrophobic residue, e.g., leucyl, isoleucyl, phenylalanyl, valyl or alanyl; (b) a cysteine or proline is substituted for (or by) another residue; (c) a residue having

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an electropositive side chain, e.g., lysyl, arginyl, or histidyl, is substituted for (or by) an electronegative residue, e.g., glutamyl or aspartyl; or (d) a residue having a bulky side chain, e.g., phenylalanine, is substituted for (or by) one not having a side chain, e.g., glycine.

Covalent modifications of prostate cancer polypeptides are included within the scope of this invention. One type of covalent modification includes reacting targeted amino acid residues of a prostate cancer polypeptide with an organic derivatizing agent that is capable of reacting with selected side chains or the N-or C-terminal residues of a prostate cancer polypeptide. Derivatization with bifunctional agents is useful, for instance, for crosslinking prostate cancer polypeptides to a water-insoluble support matrix or surface for use in the method for purifying anti-prostate cancer polypeptide antibodies or screening assays, as is more fully described below. Commonly used crosslinking agents include, e.g., 1,1-bis(diazoacetyl)-2-phenylethane, glutaraldehyde, N-hydroxysuccinimide esters, e.g., esters with 4-azidosalicylic acid, homobifunctional imidoesters, including disuccinimidyl esters such as 3,3'-dithiobis(succinimidylpropionate), bifunctional maleimides such as bis-N-maleimido-1,8-octane and agents such as methyl-3-((p-azidophenyl)dithio)propioimidate.

Other modifications include deamidation of glutaminyl and asparaginyl residues to the corresponding glutamyl and aspartyl residues, respectively, hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of serinyl, threonyl or tyrosyl residues, methylation of the amino groups of the lysine, arginine, and histidine side chains (e.g., pp. 79-86, Creighton (1983) Proteins: Structure and Molecular Properties Freeman), acetylation of the N-terminal amine, and amidation of a C-terminal carboxyl group.

Another type of covalent modification of the prostate cancer polypeptide included within the scope of this invention comprises altering the native glycosylation pattern of the polypeptide. "Altering the native glycosylation pattern" is intended for purposes herein to mean deleting one or more carbohydrate moieties found in native sequence prostate cancer polypeptide, and/or adding one or more glycosylation sites that are not present in the native sequence prostate cancer polypeptide. Glycosylation patterns can be altered in many ways. For example the use of different cell types to express prostate cancer-associated sequences can result in different glycosylation patterns.

Addition of glycosylation sites to prostate cancer polypeptides may also be accomplished by altering the amino acid sequence thereof. The alteration may be made, e.g., by the addition of, or substitution by, one or more serine or threonine residues to the native

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sequence prostate cancer polypeptide (for O-linked glycosylation sites). The prostate cancer amino acid sequence may optionally be altered through changes at the DNA level, particularly by mutating the DNA encoding the prostate cancer polypeptide at preselected bases such that codons are generated that will translate into the desired amino acids.

Another means of increasing the number of carbohydrate moieties on the prostate cancer polypeptide is by chemical or enzymatic coupling of glycosides to the polypeptide. Such methods are described in the art, e.g., in WO 87/05330, and pp. 259-306 in Aplin and Wriston (1981) CRC Crit. Rev. Biochem.

Removal of carbohydrate moieties present on the prostate cancer polypeptide may be accomplished chemically or enzymatically or by mutational substitution of codons encoding for amino acid residues that serve as targets for glycosylation. Chemical deglycosylation techniques are known in the art and described, e.g., by Hakimuddin, et al. (1987) Arch. Biochem. Biophys. 259:52-57; and Edge, et al. (1981) Anal. Biochem. 118:131-137. Enzymatic cleavage of carbohydrate moieties on polypeptides can be achieved by the use of a variety of endo-and exo-glycosidases as described by Thotakura, et al. (1987) Meth. Enzymol. 138:350-359.

Another type of covalent modification of prostate cancer comprises linking the prostate cancer polypeptide to one of a variety of nonproteinaceous polymers, e.g., polyethylene glycol, polypropylene glycol, or polyoxyalkylenes, in the manner set forth in U.S. Patent Nos. 4,640,835; 4,496,689; 4,301,144; 4,670,417; 4,791,192; or 4,179,337.

Prostate cancer polypeptides of the present invention may also be modified in a way to form chimeric molecules comprising a prostate cancer polypeptide fused to another, heterologous polypeptide or amino acid sequence. In one embodiment, such a chimeric molecule comprises a fusion of a prostate cancer polypeptide with a tag polypeptide which provides an epitope to which an anti-tag antibody can selectively bind. The epitope tag is generally placed at the amino-or carboxyl-terminus of the prostate cancer polypeptide. The presence of such epitope-tagged forms of a prostate cancer polypeptide can be detected using an antibody against the tag polypeptide. Also, provision of the epitope tag enables the prostate cancer polypeptide to be readily purified by affinity purification using an anti-tag antibody or another type of affinity matrix that binds to the epitope tag. In an alternative embodiment, the chimeric molecule may comprise a fusion of a prostate cancer polypeptide

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with an immunoglobulin or a particular region of an immunoglobulin. For a bivalent form of the chimeric molecule, such a fusion could be to the Fc region of an IgG molecule.

Various tag polypeptides and their respective antibodies are well known in the art. Examples include poly-histidine (poly-his) or poly-histidine-glycine (poly-his-gly) tags; HIS6 and metal chelation tags, the flu HA tag polypeptide and its antibody 12CA5 (Field, et al. (1988) Mol. Cell. Biol. 8:2159-2165; the c-myc tag and the 8F9, 3C7, 6E10, G4, B7, and 9E10 antibodies thereto (Evan, et al. (1985) Molecular and Cellular Biology 5:3610-3616); and the Herpes Simplex virus glycoprotein D (gD) tag and its antibody (Paborsky, et al. (1990) Protein Engineering 3:547-553). Other tag polypeptides include the Flag-peptide (Hopp, et al. (1988) BioTechnology 6:1204-1210); the KT3 epitope peptide (Martin, et al. (1992) Science 255:192-194); tubulin epitope peptide (Skinner, et al. (1991) J. Biol. Chem. 266:15163-15166); and the T7 gene 10 protein peptide tag (Lutz-Freyermuth, et al. (1990) Proc. Natl. Acad. Sci. USA 87:6393-6397).

Also included are other prostate cancer proteins of the prostate cancer family, and prostate cancer proteins from other organisms, which are cloned and expressed as outlined below. Thus, probe or degenerate polymerase chain reaction (PCR) primer sequences may be used to find other related prostate cancer proteins from humans or other organisms. As will be appreciated by those in the art, particularly useful probe and/or PCR primer sequences include the unique areas of the prostate cancer nucleic acid sequence. As is generally known in the art, preferred PCR primers are from about 15 to about 35 nucleotides in length, with from about 20 to about 30 being preferred, and may contain inosine as needed. The conditions for the PCR reaction are well known in the art (e.g., Innis, PCR Protocols, supra).

## Antibodies to prostate cancer proteins

In a preferred embodiment, when the prostate cancer protein is to be used to generate antibodies, e.g., for immunotherapy or immunodiagnosis, the prostate cancer protein should share at least one epitope or determinant with the full length protein. By "epitope" or "determinant" herein is typically meant a portion of a protein which will generate and/or bind an antibody or T-cell receptor in the context of MHC. Thus, in most instances, antibodies made to a smaller prostate cancer protein will be able to bind to the full-length protein, particularly linear epitopes. In a preferred embodiment, the epitope is unique; that is, antibodies generated to a unique epitope show little or no cross-reactivity.

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Methods of preparing polyclonal antibodies are known to the skilled artisan (e.g., Coligan, supra; and Harlow and Lane, supra). Polyclonal antibodies can be raised in a mammal, e.g., by one or more injections of an immunizing agent and, if desired, an adjuvant. Typically, the immunizing agent and/or adjuvant will be injected in the mammal by multiple subcutaneous or intraperitoneal injections. The immunizing agent may include a protein encoded by a nucleic acid of the figures or fragment thereof or a fusion protein thereof. It may be useful to conjugate the immunizing agent to a protein known to be immunogenic in the mammal being immunized. Examples of such immunogenic proteins include but are not limited to keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, and soybean trypsin inhibitor. Examples of adjuvants which may be employed include Freund's complete adjuvant and MPL-TDM adjuvant (monophosphoryl Lipid A, synthetic trehalose dicorynomycolate). The immunization protocol may be selected by one skilled in the art without undue experimentation.

The antibodies may, alternatively, be monoclonal antibodies. Monoclonal antibodies may be prepared using hybridoma methods, such as those described by Kohler and Milstein (1975) Nature 256:495. In a hybridoma method, a mouse, hamster, or other appropriate host animal, is typically immunized with an immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. Alternatively, the lymphocytes may be immunized in vitro. The immunizing agent will typically include a polypeptide encoded by a nucleic acid of Tables 1A-4 or fragment thereof, or a fusion protein thereof. Generally, either peripheral blood lymphocytes ("PBLs") are used if cells of human origin are desired, or spleen cells or lymph node cells are used if nonhuman mammalian sources are desired. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell (see pp. 59-103 in Goding (1986) Monoclonal Antibodies: Principles and Practice Academic Press). Immortalized cell lines are usually transformed mammalian cells, particularly myeloma cells of rodent, bovine and human origin. Usually, rat or mouse myeloma cell lines are employed. The hybridoma cells may be cultured in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, immortalized cells. For example, if the parental cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium

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for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine ("HAT medium"), which substances prevent the growth of HGPRT-deficient cells.

In one embodiment, the antibodies are bispecific antibodies. Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens or that have binding specificities for two epitopes on the same antigen. In one embodiment, one of the binding specificities is for a protein encoded by a nucleic acid of Tables 1A-4 or a fragment thereof, the other one is for another antigen, and preferably for a cell-surface protein or receptor or receptor subunit, preferably one that is tumor specific. Alternatively, tetramer-type technology may create multivalent reagents.

In a preferred embodiment, the antibodies to prostate cancer protein are capable of reducing or eliminating a biological function of a prostate cancer protein, as is described below. That is, the addition of anti-prostate cancer protein antibodies (either polyclonal or preferably monoclonal) to prostate cancer tissue (or cells containing prostate cancer) may reduce or eliminate the prostate cancer. Generally, at least a 25% decrease in activity, growth, size or the like is preferred, with at least about 50% being particularly preferred and about a 95-100% decrease being especially preferred.

In a preferred embodiment the antibodies to the prostate cancer proteins are humanized antibodies (e.g., Xenerex Biosciences; Medarex, Inc.; Abgenix, Inc.; Protein Design Labs, Inc.). Humanized forms of non-human (e.g., murine) antibodies are chimeric molecules of immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')2 or other antigen-binding subsequences of antibodies) which contain minimal sequence derived from non-human immunoglobulin. Humanized antibodies include human immunoglobulins (recipient antibody) in which residues from a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a nonhuman species (donor antibody) such as mouse, rat or rabbit having the desired specificity, affinity and capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies may also comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, a humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the framework (FR) regions are those of a human

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immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin (Jones, et al. (1986) Nature 321:522-525; Riechmann, et al. (1988) Nature 332:323-329; and Presta (1992) Curr. Op. Struct. Biol. 2:593-596). Humanization can be essentially performed following methods of Winter and co-workers (see, e.g., Jones, et al. (1986) Nature 321:522-525; Riechmann, et al. (1988) Nature 332:323-327; and Verhoeyen, et al. (1988) Science 239:1534-1536), by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such humanized antibodies are chimeric antibodies (U.S. Patent No. 4,816,567), wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species.

Human antibodies can also be produced using various techniques known in the art, including phage display libraries (Hoogenboom and Winter (1991) J. Mol. Biol. 227:381-388; Marks, et al. (1991) J. Mol. Biol. 222:581-597) or the preparation of human monoclonal antibodies (e.g., p77 in Cole, et al. (1985) Monoclonal Antibodies and Cancer Therapy Liss; and Boerner, et al. (1991) J. Immunol. 147(1):86-95). Similarly, human antibodies can be made by introducing of human immunoglobulin loci into transgenic animals, e.g., mice in which the endogenous immunoglobulin genes have been partially or completely inactivated. Upon challenge, human antibody production is observed, which closely resembles that seen in humans in most respects, including gene rearrangement, assembly, and antibody repertoire. This approach is described, e.g., in U.S. Patent Nos. 5,545,807; 5,545,806; 5,569,825; 5,625,126; 5,633,425; 5,661,016, and in the following scientific publications: Marks, et al. (1992) Bio/Technology 10:779-783; Lonberg, et al. (1994) Nature 368:856-859; Morrison (1994) Nature 368:812-13; Fishwild, et al. (1996) Nature Biotechnology 14:845-51; Neuberger (1996) Nature Biotechnology 14:826; Lonberg and Huszar (1995) Intern. Rev. Immunol. 13:65-93.

By immunotherapy is meant treatment of prostate cancer with an antibody raised against prostate cancer proteins. As used herein, immunotherapy can be passive or active. Passive immunotherapy as defined herein is the passive transfer of antibody to a recipient (patient). Active immunization is the induction of antibody and/or T-cell responses in a recipient (patient). Induction of an immune response is the result of providing the recipient with an antigen to which antibodies are raised. As appreciated by one of ordinary skill in the

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art, the antigen may be provided by injecting a polypeptide against which antibodies are desired to be raised into a recipient, or contacting the recipient with a nucleic acid capable of expressing the antigen and under conditions for expression of the antigen, leading to an immune response.

In a preferred embodiment the prostate cancer proteins against which antibodies are raised are secreted proteins as described above. Without being bound by theory, antibodies used for treatment, bind and prevent the secreted protein from binding to its receptor, thereby inactivating the secreted prostate cancer protein.

In another preferred embodiment, the prostate cancer protein to which antibodies are raised is a transmembrane protein. Without being bound by theory, antibodies used for treatment bind the extracellular domain of the prostate cancer protein and prevent it from binding to other proteins, such as circulating ligands or cell-associated molecules. The antibody may cause down-regulation of the transmembrane prostate cancer protein. As will be appreciated by one of ordinary skill in the art, the antibody may be a competitive, noncompetitive or uncompetitive inhibitor of protein binding to the extracellular domain of the prostate cancer protein. The antibody is also often an antagonist of the prostate cancer protein. Further, the antibody may prevent activation of the transmembrane prostate cancer protein. In one aspect, when the antibody prevents the binding of other molecules to the prostate cancer protein, the antibody prevents growth of the cell. The antibody may also be used to target or sensitize the cell to cytotoxic agents, including, but not limited to TNF-α, TNF-β, IL-1, INF-γ, and IL-2, or chemotherapeutic agents including 5FU, vinblastine, actinomycin D, cisplatin, methotrexate, and the like. In some instances the antibody belongs to a sub-type that activates serum complement when complexed with the transmembrane protein thereby mediating cytotoxicity or antigen-dependent cytotoxicity (ADCC). Thus, prostate cancer is treated by administering to a patient antibodies directed against the transmembrane prostate cancer protein. Antibody-labeling may activate a co-toxin, localize a toxin payload, or otherwise provide means to locally ablate cells.

In another preferred embodiment, the antibody is conjugated to an effector moiety. The effector moiety can be a labeling moiety such as a radioactive label or fluorescent label, or can be a therapeutic moiety. In one aspect the therapeutic moiety is a small molecule that modulates the activity of the prostate cancer protein. In another aspect the therapeutic moiety modulates the activity of molecules associated with or in close proximity to the prostate

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cancer protein. The therapeutic moiety may inhibit enzymatic activity such as protease or collagenase or protein kinase activity associated with prostate cancer.

In a preferred embodiment, the therapeutic moiety can also be a cytotoxic agent. In this method, targeting the cytotoxic agent to prostate cancer tissue or cells, results in a reduction in the number of afflicted cells, thereby reducing symptoms associated with prostate cancer. Cytotoxic agents are numerous and varied and include, but are not limited to, cytotoxic drugs or toxins or active fragments of such toxins. Suitable toxins and their corresponding fragments include diphtheria A chain, exotoxin A chain, ricin A chain, abrin A chain, curcin, crotin, phenomycin, enomycin, saporin, auristatin, and the like. Cytotoxic agents also include radiochemicals made by conjugating radioisotopes to antibodies raised against prostate cancer proteins, or binding of a radionuclide to a chelating agent that has been covalently attached to the antibody. Targeting the therapeutic moiety to transmembrane prostate cancer proteins not only serves to increase the local concentration of therapeutic moiety in the prostate cancer afflicted area, but also serves to reduce deleterious side effects, e.g., by binding to normal tissues, that may be associated with the therapeutic moiety.

In another preferred embodiment, the prostate cancer protein against which the antibodies are raised is an intracellular protein. In this case, the antibody may be conjugated to a protein which facilitates entry into the cell. In one case, the antibody enters the cell by endocytosis. In another embodiment, a nucleic acid encoding the antibody is administered to the individual or cell. Moreover, wherein the prostate cancer protein can be targeted within a cell, i.e., the nucleus, an antibody thereto contains a signal for that target localization, i.e., a nuclear localization signal.

The prostate cancer antibodies of the invention specifically bind to prostate cancer proteins. By "specifically bind" herein is meant that the antibodies bind to the protein with a  $K_d$  of at least about 0.1 mM, more usually at least about 1  $\mu$ M, preferably at least about 0.1  $\mu$ M or better, and most preferably, 0.01  $\mu$ M or better. Selectivity of binding is also important.

## Detection of prostate cancer sequence for diagnostic and therapeutic applications

In one aspect, the RNA expression levels of genes are determined for different cellular states in the prostate cancer phenotype. After androgen ablation therapy, cells that survive the therapy undergo a period of quiescence followed at sometime later by active cell

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division. As explained above, there are a variety of expression patterns characteristic of the prostate cancer genes involved in androgen-independent prostate cancer. Some genes are expressed early in the time course following ablation therapy, then drop off in expression, and then express again with emergence of androgen-independence (hi-lo-hi pattern in 1A). Other genes are expressed early in the time course following ablation therapy, then drop off in expression, and do not express again with emergence of androgen-independence (hi-lo-lo pattern in Table 1A). Still other genes are not expressed early in the time course, but express only with emergence of androgen-independence (lo-lo-hi pattern in Table 1A). Other genes are not expressed early in the time course, but then express as androgen is withdrawn and continue to express with emergence of androgen-independence (lo-hi-hi pattern in Table 1A). Finally, some genes are not expressed early in the time course, but then express as androgen is withdrawn and drop off again with emergence of androgen-independence (lo-hi-lo pattern in Table 1A). Thus, the data suggest that different antigens are expressed in quiescent cells and actively dividing androgen-independent prostate cancer cells.

In another aspect, the RNA expression levels of genes are determined for different cellular states in the prostate cancer phenotype. After androgen ablation therapy, cells that survive the therapy undergo a period of quiescence followed at sometime later by active cell division. As explained above, there are a variety of expression patterns characteristic of the prostate cancer genes involved in androgen-independent prostate cancer. Some genes are expressed early in the time course following ablation therapy, then drop off in expression, and then express again with emergence of androgen-independence (hi-lo-lo-hi pattern in Table 2A). Other genes are expressed early in the time course following ablation therapy, then drop off in expression, and do not express again with emergence of androgenindependence (hi-lo-lo-lo and hi-hi-lo-lo pattern in Table 2A). Still other genes are not expressed early in the time course, but express only with emergence of androgenindependence (lo-lo-lo-hi pattern in Table 2A). Other genes are not expressed early in the time course, but then express as androgen is withdrawn and continue to express with emergence of androgen-independence (lo-lo-hi-hi pattern in Table 2A). Finally, some genes are not expressed early in the time course, but then express as androgen is withdrawn and drop off again with emergence of androgen-independence (lo-lo-hi-lo pattern in Table 2A). Thus, the data suggest that different antigens are expressed in quiescent cells (during androgen withdrawal) and actively dividing androgen-independent prostate cancer cells.

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Effective therapy to combat androgen-independent prostate cancer requires that the timing of therapy coincide with expression of the target genes. Patients can be monitored for the expression of certain diagnostic antigens that indicate the presence of quiescent cells or which indicate the transition to actively dividing androgen-independent prostate cancer cells. Thus, therapy to combat androgen-independent prostate cancer should begin at some time following androgen ablation therapy, depending on the particular target. Typically the transition from quiescence to actively dividing androgen-independent prostate cancer occurs between 6-24 months following androgen ablation therapy. Thus, preferred time periods for the therapies of the invention are as follows:

Expression levels of genes in normal tissue (i.e., not undergoing prostate cancer) and in prostate cancer tissue (and in some cases, for varying severities of prostate cancer that relate to prognosis, as outlined below) or in non-malignant disease are evaluated to provide expression profiles. An expression profile of a particular cell state or point of development is essentially a "fingerprint" of the state. While two states may have a particular gene similarly expressed, the evaluation of a number of genes simultaneously allows the generation of a gene expression profile that is reflective of the state of the cell. By comparing expression profiles of cells in different states, information regarding which genes are important (including both up- and down-regulation of genes) in each of these states is obtained. Then, diagnosis may be performed or confirmed to determine whether a tissue sample has the gene expression profile of normal or cancerous tissue. This will provide for molecular diagnosis of related conditions.

"Differential expression," or grammatical equivalents as used herein, refers to qualitative or quantitative differences in the temporal and/or cellular gene expression patterns within and among cells and tissue. Thus, a differentially expressed gene can qualitatively have its expression altered, including an activation or inactivation, in, e.g., normal versus prostate cancer tissue. Genes may be turned on or turned off in a particular state, relative to another state thus permitting comparison of two or more states. A qualitatively regulated gene will exhibit an expression pattern within a state or cell type which is detectable by standard techniques. Some genes will be expressed in one state or cell type, but not in both. Alternatively, the difference in expression may be quantitative, e.g., in that expression is increased or decreased; i.e., gene expression is either upregulated, resulting in an increased amount of transcript, or downregulated, resulting in a decreased amount of transcript. The

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degree to which expression differs need only be large enough to quantify via standard characterization techniques as outlined below, such as by use of Affymetrix GeneChip<sup>TM</sup> expression arrays, Lockhart (1996) Nature Biotechnology 14:1675-1680, hereby expressly incorporated by reference. Other techniques include, but are not limited to, quantitative reverse transcriptase PCR, northern analysis and RNase protection. As outlined above, preferably the change in expression (i.e., upregulation or downregulation) is at least about 50%, more preferably at least about 100%, more preferably at least about 150%, more preferably at least about 200%, with from 300 to at least 1000% being especially preferred.

Evaluation may be at the gene transcript, or the protein level. The amount of gene expression may be monitored using nucleic acid probes to the DNA or RNA equivalent of the gene transcript, and the quantification of gene expression levels, or, alternatively, the final gene product itself (protein) can be monitored, e.g., with antibodies to the prostate cancer protein and standard immunoassays (ELISAs, etc.) or other techniques, including mass spectroscopy assays, 2D gel electrophoresis assays, etc. Proteins corresponding to prostate cancer genes, i.e., those identified as being important in a prostate cancer or disease phenotype, can be evaluated in a prostate cancer diagnostic test.

In a preferred embodiment, gene expression monitoring is performed simultaneously on a number of genes. Multiple protein expression monitoring can be performed as well. Similarly, these assays may be performed on an individual basis as well.

In this embodiment, the prostate cancer nucleic acid probes are attached to biochips as outlined herein for the detection and quantification of prostate cancer sequences in a particular cell. The assays are further described below in the example. PCR techniques can be used to provide greater sensitivity.

In a preferred embodiment nucleic acids encoding the prostate cancer protein are detected. Although DNA or RNA encoding the prostate cancer protein may be detected, of particular interest are methods wherein an mRNA encoding a prostate cancer protein is detected. Probes to detect mRNA can be a nucleotide/deoxynucleotide probe that is complementary to and hybridizes with the mRNA and includes, but is not limited to, oligonucleotides, cDNA, or RNA. Probes also should contain a detectable label, as defined herein. In one method the mRNA is detected after immobilizing the nucleic acid to be examined on a solid support such as nylon membranes and hybridizing the probe with the sample. Following washing to remove the non-specifically bound probe, the label is

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detected. In another method detection of the mRNA is performed in situ (in situ hybridization or ISH). In this method permeabilized cells or tissue samples are contacted with a detectably labeled nucleic acid probe for sufficient time to allow the probe to hybridize with the target mRNA. Following washing to remove the non-specifically bound probe, the label is detected. For example a digoxygenin labeled riboprobe (RNA probe) that is complementary to the mRNA encoding a prostate cancer protein is detected by binding the digoxygenin with an anti-digoxygenin secondary antibody and developed with nitro blue tetrazolium and 5-bromo-4-chloro-3-indoyl phosphate.

In a preferred embodiment, various proteins from the three classes of proteins as described herein (secreted, transmembrane, or intracellular proteins) are used in diagnostic assays. The prostate cancer proteins, antibodies, nucleic acids, modified proteins and cells containing prostate cancer sequences are used in diagnostic assays. Such may evaluate tissues, e.g., immunohistochemistry, or evaluate body fluids, e.g., blood. The detection may be direct of cells, or indirect, e.g., of products from cells. This can be performed on an individual gene or corresponding polypeptide level. In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes and/or corresponding polypeptides.

As described and defined herein, prostate cancer proteins, including intracellular, transmembrane, or secreted proteins, find use as prognostic or diagnostic markers of prostate cancer or other prostate conditions. Detection of these proteins in putative prostate cancer tissue allows for detection, diagnosis, or prognosis of prostate proliferative disorders (malignant and non-malignant) including benign prostate hyperplasia (BPH) and cancer, and prostatitis. Diagnosis may also assist in selecting a therapeutic strategy, e.g., based on expression profiles and/or comparison to archival samples. In one embodiment, antibodies are used to detect prostate cancer proteins, directly or indirectly. A preferred method separates proteins from a sample by electrophoresis on a gel (typically a denaturing and reducing protein gel, but may be another type of gel, including isoelectric focusing gels and the like). Following separation of proteins, the prostate cancer protein is detected, e.g., by immunoblotting with antibodies raised against the prostate cancer protein. Methods of immunoblotting are well known to those of ordinary skill in the art.

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In another preferred method, antibodies to the prostate cancer protein find use in in situ imaging techniques, e.g., in histology and/or in immunohistochemistry (e.g., Asai (ed. 1993) Methods in Cell Biology: Antibodies in Cell Biology (vol. 37) Academic Press. In this method cells are contacted with from one to many antibodies to the prostate cancer protein(s). Following washing to remove non-specific antibody binding, the presence of the antibody or antibodies is detected. In one embodiment the antibody is detected by incubating with a secondary antibody that contains a detectable label. In another method the primary antibody to the prostate cancer protein(s) contains a detectable label, e.g., an enzyme marker that can act on a substrate. In another preferred embodiment each one of multiple primary antibodies contains a distinct and detectable label. This method finds particular use in simultaneous screening for a plurality of prostate cancer proteins. As will be appreciated by one of ordinary skill in the art, many other histological imaging techniques are also provided by the invention.

In a preferred embodiment the label is detected in a fluorometer which has the ability to detect and distinguish emissions of different wavelengths. In addition, a fluorescence activated cell sorter (FACS) can be used in the method.

In another preferred embodiment, antibodies find use in diagnosing prostate cancer from blood, serum, plasma, stool, and other samples. Such samples, therefore, are useful as samples to be probed or tested for the presence of prostate cancer proteins, which may be diagnostic of prostate conditions beyond cancer, e.g., BPH. Antibodies can be used to detect a prostate cancer protein by previously described immunoassay techniques including ELISA, immunoblotting (western blotting), immunoprecipitation, BIACORE technology, and the like. Conversely, the presence of antibodies may indicate an immune response against an endogenous prostate cancer protein.

In a preferred embodiment, in situ hybridization of labeled prostate cancer nucleic acid probes to tissue arrays is done. For example, arrays of tissue samples, including prostate cancer tissue and/or normal tissue, are made. In situ hybridization (see, e.g., Ausubel, supra) is then performed. When comparing the fingerprints between an individual and a standard, the skilled artisan can make a diagnosis, a prognosis, or a prediction based on the findings. It is further understood that the genes which indicate the diagnosis may differ from those which indicate the prognosis and molecular profiling of the condition of the cells may lead to distinctions between responsive or refractory conditions or may be predictive of outcomes.

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In a preferred embodiment, the prostate cancer proteins, antibodies, nucleic acids, modified proteins, and cells containing prostate cancer sequences are used in prognosis assays. As above, gene expression profiles can be generated that correlate to prostate cancer or other prostate disorders, in terms of useful aspects of clinical condition, pathology, or other information which may be relevant to long term prognosis. Again, this may be done on either a protein or gene level, with the use of genes being preferred. Single or multiple genes may be useful in various combinations. As above, prostate cancer probes may be attached to biochips for the detection and quantification of prostate cancer sequences in a tissue or patient. The assays proceed as outlined above for diagnosis. PCR method may provide more sensitive and accurate quantification.

## Assays for therapeutic compounds

In a preferred embodiment members of the proteins, nucleic acids, and antibodies as described herein are used in drug screening assays. The prostate cancer proteins, antibodies, nucleic acids, modified proteins, and cells containing prostate cancer sequences are used in drug screening assays or by evaluating the effect of drug candidates on a "gene expression profile" or expression profile of polypeptides. In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes after treatment with a candidate agent (e.g., Zlokarnik, et al. (1998) Science 279:84-88; Heid (1996) Genome Res. 6:986-94).

In a preferred embodiment, the prostate cancer proteins, antibodies, nucleic acids, modified proteins, and cells containing the native or modified prostate cancer proteins are used in screening assays. That is, the present invention provides novel methods for screening for compositions which modulate the prostate cancer phenotype or an identified physiological function of a prostate cancer protein. As above, this can be done on an individual gene level or by evaluating the effect of drug candidates on a "gene expression profile". In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes after treatment with a candidate agent, see Zlokarnik, supra.

Having identified the differentially expressed genes herein, a variety of assays may be executed. In a preferred embodiment, assays may be run on an individual gene or protein level. That is, having identified a particular gene as up regulated in prostate cancer, test

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compounds can be screened for the ability to modulate gene expression or for binding to the prostate cancer protein. "Modulation" thus includes both an increase and a decrease in gene expression. The preferred amount of modulation will depend on the original change of the gene expression in normal versus tissue undergoing prostate cancer, with changes of at least 10%, preferably 50%, more preferably 100-300%, and in some embodiments 300-1000% or greater. Thus, if a gene exhibits a 4-fold increase in prostate cancer tissue compared to normal tissue, a decrease of about four-fold is often desired; similarly, a 10-fold decrease in prostate cancer tissue compared to normal tissue often provides a target value of a 10-fold increase in expression to be induced by the test compound.

The amount of gene expression may be monitored using nucleic acid probes and the quantification of gene expression levels, or, alternatively, the gene product itself can be monitored, e.g., through the use of antibodies to the prostate cancer protein and standard immunoassays. Proteomics and separation techniques may also allow quantification of expression.

In a preferred embodiment, gene expression or protein monitoring of a number of entities, i.e., an expression profile, is monitored simultaneously. Such profiles will typically involve a plurality of those entities described herein.

In this embodiment, the prostate cancer nucleic acid probes are attached to biochips as outlined herein for the detection and quantification of prostate cancer sequences in a particular cell. Alternatively, PCR may be used. Thus, a series, e.g., of microtiter plate, may be used with dispensed primers in desired wells. A PCR reaction can then be performed and analyzed for each well.

Expression monitoring can be performed to identify compounds that modify the expression of one or more prostate cancer-associated sequences, e.g., a polynucleotide sequence set out in Tables 1A-4. Generally, in a preferred embodiment, a test modulator is added to the cells prior to analysis. Moreover, screens are also provided to identify agents that modulate prostate cancer, modulate prostate cancer proteins, bind to a prostate cancer protein, or interfere with the binding of a prostate cancer protein and an antibody or other binding partner.

The term "test compound" or "drug candidate" or "modulator" or grammatical equivalents as used herein describes a molecule, e.g., protein, oligopeptide, small organic molecule, polysaccharide, polynucleotide, etc., to be tested for the capacity to directly or

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indirectly alter the prostate cancer phenotype or the expression of a prostate cancer sequence, e.g., a nucleic acid or protein sequence. In preferred embodiments, modulators alter expression profiles, or expression profile nucleic acids or proteins provided herein. In one embodiment, the modulator suppresses a prostate cancer phenotype, e.g., to a normal or non-malignant tissue fingerprint. In another embodiment, a modulator induced a prostate cancer phenotype. Generally, a plurality of assay mixtures are run in parallel with different agent concentrations to obtain a differential response to the various concentrations. Typically, one of these concentrations serves as a negative control, i.e., at zero concentration or below the level of detection.

Drug candidates encompass numerous chemical classes, though typically they are organic molecules, preferably small organic compounds having a molecular weight of more than 100 and less than about 2,500 daltons. Preferred small molecules are less than 2000, or less than 1500, or less than 1000, or less than 500 D. Candidate agents comprise functional groups necessary for structural interaction with proteins, particularly hydrogen bonding, and typically include at least an amine, carbonyl, hydroxyl or carboxyl group, preferably at least two of the functional chemical groups. The candidate agents often comprise cyclical carbon or heterocyclic structures and/or aromatic or polyaromatic structures substituted with one or more of the above functional groups. Candidate agents are also found among biomolecules including peptides, saccharides, fatty acids, steroids, purines, pyrimidines, derivatives, structural analogs, or combinations thereof. Particularly preferred are peptides.

In one aspect, a modulator will neutralize the effect of a prostate cancer protein. By "neutralize" is meant that activity of a protein is inhibited or blocked and the consequent effect on the cell.

In certain embodiments, combinatorial libraries of potential modulators will be screened for an ability to bind to a prostate cancer polypeptide or to modulate activity. Conventionally, new chemical entities with useful properties are generated by identifying a chemical compound (called a "lead compound") with some desirable property or activity, e.g., inhibiting activity, creating variants of the lead compound, and evaluating the property and activity of those variant compounds. Often, high throughput screening (HTS) methods are employed for such an analysis.

In one preferred embodiment, high throughput screening methods involve providing a library containing a large number of potential therapeutic compounds (candidate

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compounds). Such "combinatorial chemical libraries" are then screened in one or more assays to identify those library members (particular chemical species or subclasses) that display a desired characteristic activity. The compounds thus identified can serve as conventional "lead compounds" or can themselves be used as potential or actual therapeutics.

A combinatorial chemical library is a collection of diverse chemical compounds generated by either chemical synthesis or biological synthesis by combining a number of chemical "building blocks" such as reagents. For example, a linear combinatorial chemical library, such as a polypeptide (e.g., mutein) library, is formed by combining a set of chemical building blocks called amino acids in most every possible way for a given compound length (i.e., the number of amino acids in a polypeptide compound). Millions of chemical compounds can be synthesized through such combinatorial mixing of chemical building blocks. Gallop, et al. (1994) J. Med. Chem. 37:1233-1251.

Preparation and screening of combinatorial chemical libraries is well known to those of skill in the art. Such combinatorial chemical libraries include, but are not limited to. peptide libraries (see, e.g., U.S. Patent No. 5,010,175, Furka (1991) Pept. Prot. Res. 37:487-493, Houghton, et al. (1991) Nature, 354:84-88), peptoids (PCT Publication No WO 91/19735), encoded peptides (PCT Publication WO 93/20242), random bio-oligomers (PCT Publication WO 92/00091), benzodiazepines (U.S. Pat. No. 5,288,514), diversomers such as hydantoins, benzodiazepines and dipeptides (Hobbs, et al. (1993) Proc. Nat. Acad. Sci. USA 90:6909-6913), vinylogous polypeptides (Hagihara, et al. (1992) J. Amer. Chem. Soc. 114:6568-xxx), nonpeptidal peptidomimetics with a Beta-D-Glucose scaffolding (Hirschmann, et al. (1992) J. Amer. Chem. Soc. 114:9217-9218), analogous organic syntheses of small compound libraries (Chen, et al. (1994) J. Amer. Chem. Soc. 116:2661xxx), oligocarbamates (Cho, et al. (1993) Science 261:1303-1305), and/or peptidyl phosphonates (Campbell, et al. (1994) J. Org. Chem. 59:658-xxx). See, generally, Gordon, et al. (1994) J. Med. Chem. 37:1385-1401), nucleic acid libraries (see, e.g., Stratagene, Corp.), peptide nucleic acid libraries (see, e.g., U.S. Patent 5,539,083), antibody libraries (see, e.g., Vaughn, et al. (1996) Nature Biotechnology 14:309-314, and PCT/US96/10287), carbohydrate libraries (see, e.g., Liang, et al. (1996) Science 274:1520-1522, and U.S. Patent No. 5,593,853), and small organic molecule libraries (see, e.g., benzodiazepines, Baum (1993) C&EN, Jan 18, page 33; isoprenoids, U.S. Patent No. 5,569,588; thiazolidinones and metathiazanones, U.S. Patent No. 5,549,974; pyrrolidines, U.S. Patent Nos. 5,525,735 and

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5,519,134; morpholino compounds, U.S. Patent No. 5,506,337; benzodiazepines, U.S. Patent No. 5,288,514; and the like).

Devices for the preparation of combinatorial libraries are commercially available (see, e.g., 357 MPS, 390 MPS, Advanced Chem Tech, Louisville KY, Symphony, Rainin, Woburn, MA, 433A Applied Biosystems, Foster City, CA, 9050 Plus, Millipore, Bedford, MA).

A number of well known robotic systems have also been developed for solution phase chemistries. These systems include automated workstations like the automated synthesis apparatus developed by Takeda Chemical Industries, LTD. (Osaka, Japan) and many robotic systems utilizing robotic arms (Zymate II, Zymark Corporation, Hopkinton, Mass.; Orca, Hewlett-Packard, Palo Alto, Calif.), which mimic the manual synthetic operations performed by a chemist. Many of the above devices are suitable for use with the present invention. The nature and implementation of modifications to these devices (if any) so that they can operate as discussed herein will be apparent to persons skilled in the relevant art. In addition, numerous combinatorial libraries are themselves commercially available (see, e.g., ComGenex, Princeton, N.J., Asinex, Moscow, Ru, Tripos, Inc., St. Louis, MO, ChemStar, Ltd, Moscow, RU, 3D Pharmaceuticals, Exton, PA, Martek Biosciences, Columbia, MD, etc.).

The assays to identify modulators are amenable to high throughput screening.

Preferred assays thus detect enhancement or inhibition of prostate cancer gene transcription, inhibition or enhancement of polypeptide expression, and inhibition or enhancement of polypeptide activity.

High throughput assays for the presence, absence, quantification, or other properties of particular nucleic acids or protein products are well known to those of skill in the art. Similarly, binding assays and reporter gene assays are similarly well known. Thus, e.g., U.S. Patent No. 5,559,410 discloses high throughput screening methods for proteins, U.S. Patent No. 5,585,639 discloses high throughput screening methods for nucleic acid binding (i.e., in arrays), while U.S. Patent Nos. 5,576,220 and 5,541,061 disclose high throughput methods of screening for ligand/antibody binding.

In addition, high throughput screening systems are commercially available (see, e.g., Zymark Corp., Hopkinton, MA; Air Technical Industries, Mentor, OH; Beckman Instruments, Inc. Fullerton, CA; Precision Systems, Inc., Natick, MA, etc.). These systems

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typically automate entire procedures, including sample and reagent pipetting, liquid dispensing, timed incubations, and final readings of the microplate in detector(s) appropriate for the assay. These configurable systems provide high throughput and rapid start up as well as a high degree of flexibility and customization. The manufacturers of such systems provide detailed protocols for various high throughput systems. Thus, e.g., Zymark Corp. provides technical bulletins describing screening systems for detecting the modulation of gene transcription, ligand binding, and the like.

In one embodiment, modulators are proteins, often naturally occurring proteins or fragments of naturally occurring proteins. Thus, e.g., cellular extracts containing proteins, or random or directed digests of proteinaceous cellular extracts, may be used. In this way libraries of proteins may be made for screening in the methods of the invention. Particularly preferred in this embodiment are libraries of bacterial, fungal, viral, and mammalian proteins, with the latter being preferred, and human proteins being especially preferred. Particularly useful test compound will be directed to the class of proteins to which the target belongs, e.g., substrates for enzymes or ligands and receptors.

In a preferred embodiment, modulators are peptides of from about 5 to about 30 amino acids, with from about 5 to about 20 amino acids being preferred, and from about 7 to about 15 being particularly preferred. The peptides may be digests of naturally occurring proteins as is outlined above, random peptides, or "biased" random peptides. By "randomized" or grammatical equivalents herein is meant that each nucleic acid and peptide consists of essentially random nucleotides and amino acids, respectively. Since generally these random peptides (or nucleic acids, discussed below) are chemically synthesized, they may typically incorporate any nucleotide or amino acid at any position. The synthetic process can be designed to generate randomized proteins or nucleic acids, to allow the formation of all or most of the possible combinations over the length of the sequence, thus forming a library of randomized candidate bioactive proteinaceous agents.

In one embodiment, the library is fully randomized, with no sequence preferences or constants at any position. In a preferred embodiment, the library is biased. That is, some positions within the sequence are either held constant, or are selected from a limited number of possibilities. For example, in a preferred embodiment, the nucleotides or amino acid residues are randomized within a defined class, e.g., of hydrophobic amino acids, hydrophilic residues, sterically biased (either small or large) residues, towards the creation of nucleic acid

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binding domains, the creation of cysteines, for cross-linking, prolines for SH-3 domains, serines, threonines, tyrosines, or histidines for phosphorylation sites, etc., or to purines, etc.

Modulators of prostate cancer can also be nucleic acids, as defined above.

As described above generally for proteins, nucleic acid modulating agents may be naturally occurring nucleic acids, random nucleic acids, or "biased" random nucleic acids. For example, digests of prokaryotic or eukaryotic genomes may be used as is outlined above for proteins.

In a preferred embodiment, the candidate compounds are organic chemical moieties, a wide variety of which are available in the literature.

After the candidate agent has been added and the cells allowed to incubate for some period of time, the sample containing a target sequence to be analyzed is added to the biochip. If required, the target sequence is prepared using known techniques. For example, the sample may be treated to lyse the cells, using known lysis buffers, electroporation, etc., with purification and/or amplification such as PCR performed as appropriate. For example, an in vitro transcription with labels covalently attached to the nucleotides is performed. Generally, the nucleic acids are labeled with biotin-FITC or PE, or with cy3 or cy5.

In a preferred embodiment, the target sequence is labeled with, e.g., a fluorescent, a chemiluminescent, a chemical, or a radioactive signal, to provide a means of detecting the target sequence's specific binding to a probe. The label also can be an enzyme, such as, alkaline phosphatase or horseradish peroxidase, which when provided with an appropriate substrate produces a product that can be detected. Alternatively, the label can be a labeled compound or small molecule, such as an enzyme inhibitor, that binds but is not catalyzed or altered by the enzyme. The label also can be a moiety or compound, such as, an epitope tag or biotin which specifically binds to streptavidin. For the example of biotin, the streptavidin is labeled as described above, thereby, providing a detectable signal for the bound target sequence. Unbound labeled streptavidin is typically removed prior to analysis.

As will be appreciated by those in the art, these assays can be direct hybridization assays or can comprise "sandwich assays", which include the use of multiple probes, as is generally outlined in U.S. Patent Nos. 5,681,702, 5,597,909, 5,545,730, 5,594,117, 5,591,584, 5,571,670, 5,580,731, 5,571,670, 5,591,584, 5,624,802, 5,635,352, 5,594,118, 5,359,100, 5,124,246, and 5,681,697, each of which is hereby incorporated by reference. In this embodiment, in general, the target nucleic acid is prepared as outlined above, and then

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added to the biochip comprising a plurality of nucleic acid probes, under conditions that allow the formation of a hybridization complex.

A variety of hybridization conditions may be used in the present invention, including high, moderate, and low stringency conditions as outlined above. The assays are generally run under stringency conditions which allows formation of the label probe hybridization complex only in the presence of target. Stringency can be controlled by altering a step parameter that is a thermodynamic variable, including, but not limited to, temperature, formamide concentration, salt concentration, chaotropic salt concentration pH, organic solvent concentration, etc.

These parameters may also be used to control non-specific binding, as is generally outlined in U.S. Patent No. 5,681,697. Thus it may be desirable to perform certain steps at higher stringency conditions to reduce non-specific binding.

The reactions outlined herein may be accomplished in a variety of ways. Components of the reaction may be added simultaneously, or sequentially, in different orders, with preferred embodiments outlined below. In addition, the reaction may include a variety of other reagents. These include salts, buffers, neutral proteins, e.g., albumin, detergents, etc., which may be used to facilitate optimal hybridization and detection, and/or reduce non-specific or background interactions. Reagents that otherwise improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc., may also be used as appropriate, depending on the sample preparation methods and purity of the target.

The assay data are analyzed to determine the expression levels, and changes in expression levels as between states, of individual genes, forming a gene expression profile.

Screens are performed to identify modulators of the prostate cancer or related phenotype. In one embodiment, screening is performed to identify modulators that can induce or suppress a particular expression profile, thus preferably generating the associated phenotype. In another embodiment, e.g., for diagnostic applications, having identified differentially expressed genes important in a particular state, screens can be performed to identify modulators that alter expression of individual genes. In an another embodiment, screening is performed to identify modulators that alter a biological function of the expression product of a differentially expressed gene. Again, having identified the importance of a gene in a particular state, screens are performed to identify agents that bind and/or modulate the biological activity of the gene product.

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In addition screens can be done for genes that are induced in response to a candidate agent. After identifying a modulator based upon its ability to suppress a prostate cancer expression pattern leading to a normal expression pattern, or to modulate a single prostate cancer gene expression profile so as to mimic the expression of the gene from normal tissue, a screen as described above can be performed to identify genes that are specifically modulated in response to the agent. Comparing expression profiles between normal tissue and agent treated prostate cancer tissue reveals genes that are not expressed in normal tissue or prostate cancer tissue, but are expressed in agent treated tissue. These agent-specific sequences can be identified and used by methods described herein for prostate cancer genes or proteins. In particular these sequences and the proteins they encode find use in marking or identifying agent treated cells. In addition, antibodies can be raised against the agent induced proteins and used to target novel therapeutics to the treated prostate cancer tissue sample.

Thus, in one embodiment, a test compound is administered to a population of prostate cancer cells, that have an associated prostate cancer expression profile. By "administration" or "contacting" herein is meant that the candidate agent is added to the cells in such a manner as to allow the agent to act upon the cell, whether by uptake and intracellular action, or by action at the cell surface. In some embodiments, nucleic acid encoding a proteinaceous candidate agent (e.g., a peptide) may be put into a viral construct such as an adenoviral or retroviral construct, and added to the cell, such that expression of the peptide agent is accomplished, e.g., PCT US97/01019. Regulatable gene therapy systems can also be used.

Once the test compound has been administered to the cells, the cells can be washed if desired and are allowed to incubate under preferably physiological conditions for some period of time. The cells are then harvested and a new gene expression profile is generated, as outlined herein.

Thus, e.g., prostate cancer or non-malignant tissue may be screened for agents that modulate, e.g., induce or suppress the prostate cancer or related phenotype. A change in at least one gene, preferably many, of the expression profile indicates that the agent has an effect on prostate cancer activity. By defining such a signature for the prostate cancer phenotype, screens for new drugs that alter the phenotype can be devised. With this approach, the drug target need not be known and need not be represented in the original expression screening platform, nor does the level of transcript for the target protein need to change.

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In a preferred embodiment, as outlined above, screens may be done on individual genes and gene products (proteins). That is, having identified a particular differentially expressed gene as important in a particular state, screening of modulators of either the expression of the gene or the gene product itself can be done. The gene products of differentially expressed genes are sometimes referred to herein as "prostate cancer proteins" or a "prostate cancer modulatory protein". The prostate cancer modulatory protein may be a fragment, or alternatively, be the full length protein to the fragment encoded by the nucleic acids of the Tables 1A-4. Preferably, the prostate cancer modulatory protein is a fragment. In a preferred embodiment, the prostate cancer amino acid sequence which is used to determine sequence identity or similarity is encoded by a nucleic acid of Tables 1A-4. In another embodiment, the sequences are naturally occurring allelic variants of a protein encoded by a nucleic acid of Tables 1A-4. In another embodiment, the sequences are sequence variants as further described herein.

Preferably, the prostate cancer modulatory protein is a fragment of approximately 14 to 24 amino acids long. More preferably the fragment is a soluble fragment. Preferably, the fragment includes a non-transmembrane region. In a preferred embodiment, the fragment has an N-terminal Cys to aid in solubility. In one embodiment, the C-terminus of the fragment is kept as a free acid and the N-terminus is a free amine to aid in coupling, i.e., to cysteine.

In one embodiment the prostate cancer proteins are conjugated to an immunogenic agent as discussed herein. In one embodiment the prostate cancer protein is conjugated to BSA.

Measurements of prostate cancer polypeptide activity, or of prostate cancer or the prostate cancer phenotype can be performed using a variety of assays. For example, the effects of the test compounds upon the function of the prostate cancer polypeptides can be measured by examining parameters described above. A suitable physiological change that affects activity can be used to assess the influence of a test compound on the polypeptides of this invention. When the functional consequences are determined using intact cells or animals, one can also measure a variety of effects such as, in the case of prostate cancer associated with tumors, tumor growth, tumor metastasis, neovascularization, hormone release, transcriptional changes to both known and uncharacterized genetic markers (e.g., northern blots), changes in cell metabolism such as cell growth or pH changes, and changes

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in intracellular second messengers such as cGMP. In the assays of the invention, a mammalian prostate cancer polypeptide is typically used, e.g., mouse, preferably human.

Assays to identify compounds with modulating activity can be performed in vitro. For example, a prostate cancer polypeptide is first contacted with a potential modulator and incubated for a suitable amount of time, e.g., from 0.5 to 48 hours. In one embodiment, the prostate cancer polypeptide levels are determined in vitro by measuring the level of protein or mRNA. The level of protein is measured using immunoassays such as western blotting, ELISA, and the like with an antibody that selectively binds to the prostate cancer polypeptide or a fragment thereof. For measurement of mRNA, amplification, e.g., using PCR, LCR, or hybridization assays, e.g., northern hybridization, RNAse protection, dot blotting, are preferred. The level of protein or mRNA is detected using directly or indirectly labeled detection agents, e.g., fluorescently or radioactively labeled nucleic acids, radioactively or enzymatically labeled antibodies, and the like, as described herein.

Alternatively, a reporter gene system can be devised using the prostate cancer protein promoter operably linked to a reporter gene such as luciferase, green fluorescent protein, CAT, or  $\beta$ -gal. The reporter construct is typically transfected into a cell. After treatment with a potential modulator, the amount of reporter gene transcription, translation, or activity is measured according to standard techniques known to those of skill in the art.

In a preferred embodiment, as outlined above, screens may be done on individual genes and gene products (proteins). That is, having identified a particular differentially expressed gene as important in a particular state, screening of modulators of the expression of the gene or the gene product itself can be done. The gene products of differentially expressed genes are sometimes referred to herein as "prostate cancer proteins." The prostate cancer protein may be a fragment, or alternatively, be the full length protein corresponding to a fragment shown herein.

In one embodiment, screening for modulators of expression of specific genes is performed. Typically, the expression of only one or a few genes are evaluated. In another embodiment, screens are designed to first find compounds that bind to differentially expressed proteins. These compounds are then evaluated for the ability to modulate differentially expressed activity. Moreover, once initial candidate compounds are identified, variants can be further screened to better evaluate structure activity relationships.

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In a preferred embodiment, binding assays are done. In general, purified or isolated gene product is used; that is, the gene products of one or more differentially expressed nucleic acids are made. For example, antibodies are generated to the protein gene products, and standard immunoassays are run to determine the amount of protein present.

Alternatively, cells comprising the prostate cancer proteins can be used in the assays.

Thus, in a preferred embodiment, the methods comprise combining a prostate cancer protein and a candidate compound, and determining the binding of the compound to the prostate cancer protein. Preferred embodiments utilize the human prostate cancer protein, although other mammalian proteins may also be used, e.g., for the development of animal models of human disease. In some embodiments, as outlined herein, variant or derivative prostate cancer proteins may be used.

Generally, in a preferred embodiment of the methods herein, the prostate cancer protein or the candidate agent is non-diffusably bound to an insoluble support having isolated sample receiving areas (e.g., a microtiter plate, an array, etc.). The insoluble supports may be made of a composition to which the compositions can be bound, is readily separated from soluble material, and is otherwise compatible with the overall method of screening. The surface of such supports may be solid or porous and of a convenient shape. Examples of suitable insoluble supports include microtiter plates, arrays, membranes, and beads. These are typically made of glass, plastic (e.g., polystyrene), polysaccharides, nylon or nitrocellulose, teflon<sup>TM</sup>, etc. Microtiter plates and arrays are especially convenient because a large number of assays can be carried out simultaneously, using small amounts of reagents and samples. The particular manner of binding of the composition should be compatible with the reagents and overall methods of the invention, maintain the activity of the composition, and be nondiffusable. Preferred methods of binding include the use of antibodies (which do not sterically block either the ligand binding site or activation sequence when the protein is bound to the support), direct binding to "sticky" or ionic supports, chemical crosslinking, the synthesis of the protein or agent on the surface, etc. Following binding of the protein or agent, excess unbound material is removed by washing. The sample receiving areas may then be blocked through incubation with bovine serum albumin (BSA), casein, or other innocuous protein or other moiety.

In a preferred embodiment, the prostate cancer protein is bound to the support, and a test compound is added to the assay. Alternatively, the candidate agent is bound to the

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support and the prostate cancer protein is added. Novel binding agents include specific antibodies, non-natural binding agents identified in screens of chemical libraries, peptide analogs, etc. Of particular interest are screening assays for agents that have a low toxicity for human cells. A wide variety of assays may be used for this purpose, including labeled in vitro protein-protein binding assays, electrophoretic mobility shift assays, immunoassays for protein binding, functional assays (phosphorylation assays, etc.) and the like.

The determination of the binding of the test modulating compound to the prostate cancer protein may be done in a number of ways. In a preferred embodiment, the compound is labeled, and binding determined directly, e.g., by attaching all or a portion of the prostate cancer protein to a solid support, adding a labeled candidate agent (e.g., a fluorescent label), washing off excess reagent, and determining whether the label is present on the solid support. Various blocking and washing steps may be utilized as appropriate.

In some embodiments, only one of the components is labeled, e.g., the proteins (or proteinaceous candidate compounds) can be labeled. Alternatively, more than one component can be labeled with different labels, e.g., <sup>125</sup>I for the proteins and a fluorophor for the compound. Proximity reagents, e.g., quenching or energy transfer reagents are also useful.

In one embodiment, the binding of the test compound is determined by competitive binding assay. The competitor is a binding moiety known to bind to the target molecule (i.e., a prostate cancer protein), such as an antibody, peptide, binding partner, ligand, etc. Under certain circumstances, there may be competitive binding between the compound and the binding moiety, with the binding moiety displacing the compound. In one embodiment, the test compound is labeled. Either the compound, or the competitor, or both, is added first to the protein for a time sufficient to allow binding, if present. Incubations may be performed at a temperature which facilitates optimal activity, typically between 4 and 40° C. Incubation periods are typically optimized, e.g., to facilitate rapid high throughput screening. Typically between 0.1 and 1 hour will be sufficient. Excess reagent is generally removed or washed away. The second component is then added, and the presence or absence of the labeled component is followed, to indicate binding.

In a preferred embodiment, the competitor is added first, followed by the test compound. Displacement of the competitor is an indication that the test compound is binding to the prostate cancer protein and thus is capable of binding to, and potentially modulating,

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the activity of the prostate cancer protein. In this embodiment, either component can be labeled. Thus, e.g., if the competitor is labeled, the presence of label in the wash solution indicates displacement by the agent. Alternatively, if the test compound is labeled, the presence of the label on the support indicates displacement.

In an alternative embodiment, the test compound is added first, with incubation and washing, followed by the competitor. The absence of binding by the competitor may indicate that the test compound is bound to the prostate cancer protein with a higher affinity. Thus, if the test compound is labeled, the presence of the label on the support, coupled with a lack of competitor binding, may indicate that the test compound is capable of binding to the prostate cancer protein.

In a preferred embodiment, the methods comprise differential screening to identity agents that are capable of modulating the activity of the prostate cancer proteins. In this embodiment, the methods comprise combining a prostate cancer protein and a competitor in a first sample. A second sample comprises a test compound, a prostate cancer protein, and a competitor. The binding of the competitor is determined for both samples, and a change, or difference in binding between the two samples indicates the presence of an agent capable of binding to the prostate cancer protein and potentially modulating its activity. That is, if the binding of the competitor is different in the second sample relative to the first sample, the agent is capable of binding to the prostate cancer protein.

Alternatively, differential screening is used to identify drug candidates that bind to the native prostate cancer protein, but cannot bind to modified prostate cancer proteins. The structure of the prostate cancer protein may be modeled, and used in rational drug design to synthesize agents that interact with that site. Drug candidates that affect the activity of a prostate cancer protein are also identified by screening drugs for the ability to either enhance or reduce the activity of the protein.

Positive controls and negative controls may be used in the assays. Preferably control and test samples are performed in at least triplicate to obtain statistically significant results. Incubation of samples is for a time sufficient for the binding of the agent to the protein. Following incubation, samples are washed free of non-specifically bound material and the amount of bound, generally labeled agent determined. For example, where a radiolabel is employed, the samples may be counted in a scintillation counter to determine the amount of bound compound.

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A variety of other reagents may be included in the screening assays. These include reagents like salts, neutral proteins, e.g., albumin, detergents, etc., which may be used to facilitate optimal protein-protein binding and/or reduce non-specific or background interactions. Also reagents that otherwise improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc., may be used. The mixture of components may be added in an order that provides for the requisite binding.

In a preferred embodiment, the invention provides methods for screening for a compound capable of modulating the activity of a prostate cancer protein. The methods comprise adding a test compound, as defined above, to a cell comprising prostate cancer proteins. Preferred cell types include almost any cell. The cells contain a recombinant nucleic acid that encodes a prostate cancer protein. In a preferred embodiment, a library of candidate agents are tested on a plurality of cells.

In one aspect, the assays are evaluated in the presence or absence or previous or subsequent exposure of physiological signals, e.g., hormones, antibodies, peptides, antigens, cytokines, growth factors, action potentials, pharmacological agents including chemotherapeutics, radiation, carcinogenics, or other cells (e.g., cell-cell contacts). In another example, the determinations are determined at different stages of the cell cycle process.

In this way, compounds that modulate prostate cancer agents are identified.

Compounds with pharmacological activity are able to enhance or interfere with the activity of the prostate cancer protein. Once identified, similar structures are evaluated to identify critical structural feature of the compound.

In one embodiment, a method of inhibiting prostate cancer cell division is provided. The method comprises administration of a prostate cancer inhibitor. In another embodiment, a method of inhibiting prostate cancer or other prostate proliferative condition is provided. The method comprises administration of a prostate cancer inhibitor. In a further embodiment, methods of treating cells or individuals with prostate cancer are provided. The method comprises administration of a prostate cancer inhibitor.

In one embodiment, a prostate cancer inhibitor is an antibody as discussed above. In another embodiment, the prostate cancer inhibitor is an antisense molecule.

A variety of cell growth, proliferation, and metastasis assays are known to those of skill in the art, as described below.

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# Soft agar growth or colony formation in suspension

Normal cells require a solid substrate to attach and grow. When the cells are transformed, they lose this phenotype and grow detached from the substrate. For example, transformed cells can grow in stirred suspension culture or suspended in semi-solid media, such as semi-solid or soft agar. The transformed cells, when transfected with tumor suppressor genes, regenerate normal phenotype and require a solid substrate to attach and grow. Soft agar growth or colony formation in suspension assays can be used to identify modulators of prostate cancer sequences, which when expressed in host cells, inhibit abnormal cellular proliferation and transformation. A therapeutic compound would reduce or eliminate the host cells' ability to grow in stirred suspension culture or suspended in semi-solid media, such as semi-solid or soft.

Techniques for soft agar growth or colony formation in suspension assays are described in Freshney (1994) <u>Culture of Animal Cells a Manual of Basic Technique</u> 3d ed. Wiley-Liss, herein incorporated by reference. See also, the methods section of Garkavtsev, et al. (1996), supra, herein incorporated by reference.

### Contact inhibition and density limitation of growth

Normal cells typically grow in a flat and organized pattern in a petri dish until they touch other cells. When the cells touch one another, they are contact inhibited and stop growing. When cells are transformed, however, the cells are not contact inhibited and continue to grow to high densities in disorganized foci. Thus, the transformed cells grow to a higher saturation density than normal cells. This can be detected morphologically by the formation of a disoriented monolayer of cells or rounded cells in foci within the regular pattern of normal surrounding cells. Alternatively, labeling index with (<sup>3</sup>H)-thymidine at saturation density can be used to measure density limitation of growth. See Freshney (1994), supra. The transformed cells, when transfected with tumor suppressor genes, regenerate a normal phenotype and become contact inhibited and would grow to a lower density.

In this assay, labeling index with (<sup>3</sup>H)-thymidine at saturation density is a preferred method of measuring density limitation of growth. Transformed host cells are transfected with a prostate cancer-associated sequence and are grown for 24 hours at saturation density in non-limiting medium conditions. The percentage of cells labeling with (<sup>3</sup>H)-thymidine is determined autoradiographically. See, Freshney (1994), supra.

#### Growth factor or serum dependence

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Transformed cells have a lower serum dependence than their normal counterparts (see, e.g., Temin (1966) <u>J. Natl. Cancer Insti.</u> 37:167-175; Eagle, et al. (1970) <u>J. Exp. Med.</u> 131:836-879); Freshney, supra. This is in part due to release of various growth factors by the transformed cells. Growth factor or serum dependence of transformed host cells can be compared with that of control.

# Tumor specific markers levels

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Tumor cells release an increased amount of certain factors (hereinafter "tumor specific markers") than their normal counterparts. For example, plasminogen activator (PA) is released from human glioma at a higher level than from normal brain cells (see, e.g., Gullino, "Angiogenesis, tumor vascularization, and potential interference with tumor growth" pp. 178-184 in Mihich (ed. 1985) <u>Biological Responses in Cancer Plenum</u>. Similarly, Tumor angiogenesis factor (TAF) is released at a higher level in tumor cells than their normal counterparts. See, e.g., Folkman (1992) <u>Angiogenesis and Cancer</u>, Sem. Cancer Biol.

Various techniques which measure the release of these factors are described in Freshney (1994), supra. Also, see, Unkless, et al. (1974) <u>J. Biol. Chem.</u> 249:4295-4305; Strickland and Beers (1976) <u>J. Biol. Chem.</u> 251:5694-5702; Whur, et al. (1980) <u>Br. J. Cancer</u> 42:305-312; Gullino, "Angiogenesis, tumor vascularization, and potential interference with tumor growth" pp. 178-184 in Mihich (ed. 1985) <u>Biological Responses in Cancer</u> Plenum; and Freshney (1985) <u>Anticancer Res.</u> 5:111-130.

### 20 Invasiveness into Matrigel

The degree of invasiveness into Matrigel or some other extracellular matrix constituent can be used as an assay to identify compounds that modulate prostate cancer-associated sequences. Tumor cells exhibit a good correlation between malignancy and invasiveness of cells into Matrigel or some other extracellular matrix constituent. In this assay, tumorigenic cells are typically used as host cells. Expression of a tumor suppressor gene in these host cells would decrease invasiveness of the host cells.

Techniques described in Freshney (1994), supra, can be used. Briefly, the level of invasion of host cells can be measured by using filters coated with Matrigel or some other extracellular matrix constituent. Penetration into the gel, or through to the distal side of the filter, is rated as invasiveness, and rated histologically by number of cells and distance moved, or by prelabeling the cells with <sup>125</sup>I and counting the radioactivity on the distal side of the filter or bottom of the dish. See, e.g., Freshney (1984), supra.

#### Tumor growth in vivo

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Effects of prostate cancer-associated sequences on cell growth can be tested in transgenic or immune-suppressed mice. Knock-out transgenic mice can be made, in which the prostate cancer gene is disrupted or in which a prostate cancer gene is inserted. Knock-out transgenic mice can be made by insertion of a marker gene or other heterologous gene into the endogenous prostate cancer gene site in the mouse genome via homologous recombination. Such mice can also be made by substituting the endogenous prostate cancer gene with a mutated version of the prostate cancer gene, or by mutating the endogenous prostate cancer gene, e.g., by exposure to carcinogens.

A DNA construct is introduced into the nuclei of embryonic stem cells. Cells containing the newly engineered genetic lesion are injected into a host mouse embryo, which is re-implanted into a recipient female. Some of these embryos develop into chimeric mice that possess germ cells partially derived from the mutant cell line. Therefore, by breeding the chimeric mice it is possible to obtain a new line of mice containing the introduced genetic lesion (see, e.g., Capecchi, et al. (1989) <a href="Science 244:1288-1292">Science 244:1288-1292</a>). Chimeric targeted mice can be derived according to Hogan, et al. (1988) <a href="Manipulating the Mouse Embryo: A Laboratory Manual CSH Press">Manual CSH Press</a>; and Robertson (ed. 1987) <a href="Teratocarcinomas and Embryonic Stem Cells: A Practical Approach IRL Press">IRL Press</a>, Washington, D.C.

Alternatively, various immune-suppressed or immune-deficient host animals can be used. For example, genetically athymic "nude" mouse (see, e.g., Giovanella, et al. (1974) <u>J. Natl. Cancer Inst.</u> 52:921-930), a SCID mouse, a thymectomized mouse, or an irradiated mouse (see, e.g., Bradley, et al. (1978) <u>Br. J. Cancer</u> 38:263-272; Selby, et al. (1980) <u>Br. J. Cancer</u> 41:52-61) can be used as a host. Transplantable tumor cells (typically about 10<sup>6</sup> cells) injected into isogenic hosts will produce invasive tumors in a high proportions of cases, while normal cells of similar origin will not. In hosts which developed invasive tumors, cells expressing a prostate cancer-associated sequences are injected subcutaneously. After a suitable length of time, preferably 4-8 weeks, tumor growth is measured (e.g., by volume or by its two largest dimensions) and compared to the control. Tumors that have statistically significant reduction (using, e.g., Student's T test) are said to have inhibited growth.

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Polynucleotide modulators of prostate cancer

<u>Antisense and RNAi Polynucleotides</u>

In certain embodiments, the activity of a prostate cancer-associated protein is down-regulated, or entirely inhibited, by the use of antisense polynucleotide, i.e., a nucleic acid complementary to, and which can preferably hybridize specifically to, a coding mRNA nucleic acid sequence, e.g., a prostate cancer protein mRNA, or a subsequence thereof. Binding of the antisense polynucleotide to the mRNA reduces the translation and/or stability of the mRNA.

In the context of this invention, antisense polynucleotides can comprise naturally-occurring nucleotides, or synthetic species formed from naturally-occurring subunits or their close homologs. Antisense polynucleotides may also have altered sugar moieties or intersugar linkages. Exemplary among these are the phosphorothioate and other sulfur containing species which are known for use in the art. Analogs are comprehended by this invention so long as they function effectively to hybridize with the prostate cancer protein mRNA. See, e.g., Isis Pharmaceuticals, Carlsbad, CA; Sequitor, Inc., Natick, MA.

Such antisense polynucleotides can readily be synthesized using recombinant means, or can be synthesized in vitro. Equipment for such synthesis is sold by several vendors, including Applied Biosystems. The preparation of other oligonucleotides such as phosphorothioates and alkylated derivatives is also well known to those of skill in the art.

Antisense molecules as used herein include antisense or sense oligonucleotides. Sense oligonucleotides can, e.g., be employed to block transcription by binding to the antisense strand. The antisense and sense oligonucleotide comprise a single-stranded nucleic acid sequence (either RNA or DNA) capable of binding to target mRNA (sense) or DNA (antisense) sequences for prostate cancer molecules. A preferred antisense molecule is for a prostate cancer sequences in Tables 1A-4, or for a ligand or activator thereof. Antisense or sense oligonucleotides, according to the present invention, comprise a fragment generally at least about 14 nucleotides, preferably from about 14 to 30 nucleotides. The ability to derive an antisense or a sense oligonucleotide, based upon a cDNA sequence encoding a given protein is described in, e.g., Stein and Cohen (1988) Cancer Res. 48:2659-2668; and van der Krol, et al. (1988) BioTechniques 6:958-976.

RNA interference is a mechanism to suppress gene expression in a sequence specific manner. See, e.g., Brumelkamp, et al. (2002) <u>Sciencexpress</u> (21March2002); Sharp (1999) <u>Genes Dev.</u> 13:139-141; and Cathew (2001) <u>Curr. Op. Cell Biol.</u> 13:244-248. In mammalian cells, short, e.g., 21 nt, double stranded small interfering RNAs (siRNA) have been shown to

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be effective at inducing an RNAi response. See, e.g., Elbashir, et al. (2001) <u>Nature</u> 411:494-498. The mechanism may be used to downregulate expression levels of identified genes, e.g., treatment of or validation of relevance to disease.

### 5 Ribozymes

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In addition to antisense polynucleotides, ribozymes can be used to target and inhibit transcription of prostate cancer-associated nucleotide sequences. A ribozyme is an RNA molecule that catalytically cleaves other RNA molecules. Different kinds of ribozymes have been described, including group I ribozymes, hammerhead ribozymes, hairpin ribozymes, RNase P, and axhead ribozymes (see, e.g., Castanotto, et al. (1994) Adv. in Pharmacology 25: 289-317 for a general review of the properties of different ribozymes).

The general features of hairpin ribozymes are described, e.g., in Hampel, et al. (1990) Nucl. Acids Res. 18:299-304; European Patent Publication No. 0 360 257; U.S. Patent No. 5,254,678. Methods of preparing are well known to those of skill in the art. See, e.g., WO 94/26877; Ojwang, et al. (1993) Proc. Natl. Acad. Sci. USA 90:6340-6344; Yamada, et al. (1994) Human Gene Therapy 1:39-45; Leavitt, et al. (1995) Proc. Natl. Acad. Sci. USA 92:699-703; Leavitt, et al. (1994) Human Gene Therapy 5:1151-120; and Yamada, et al. (1994) Virology 205:121-126.

Polynucleotide modulators of prostate cancer may be introduced into a cell containing the target nucleotide sequence by formation of a conjugate with a ligand binding molecule, as described in WO 91/04753. Suitable ligand binding molecules include, but are not limited to, cell surface receptors, growth factors, other cytokines, or other ligands that bind to cell surface receptors. Preferably, conjugation of the ligand binding molecule does not substantially interfere with the ability of the ligand binding molecule to bind to its corresponding molecule or receptor, or block entry of the sense or antisense oligonucleotide or its conjugated version into the cell. Alternatively, a polynucleotide modulator of prostate cancer may be introduced into a cell containing the target nucleic acid sequence, e.g., by formation of an polynucleotide-lipid complex, as described in WO 90/10448. It is understood that the use of antisense molecules or knock out and knock in models may also be used in screening assays as discussed above, in addition to methods of treatment.

Thus, in one embodiment, methods of modulating prostate disorders, e.g., cancer in cells or organisms, are provided. In one embodiment, the methods comprise administering to

a patient, e.g., to a cell within the patient, an anti-prostate cancer antibody that reduces or eliminates the biological activity of an endogenous prostate cancer protein. Alternatively, the methods comprise administering to a cell or organism a recombinant nucleic acid encoding a prostate cancer protein. This may be accomplished in many ways. In a preferred embodiment, e.g., when the prostate cancer sequence is down-regulated in prostate cancer, such state may be reversed by increasing the amount of prostate cancer gene product in the cell. This can be accomplished, e.g., by overexpressing the endogenous prostate cancer gene or administering a gene encoding the prostate cancer sequence, using known gene-therapy techniques, e.g.. In a preferred embodiment, the gene therapy techniques include the incorporation of the exogenous gene using enhanced homologous recombination (EHR), e.g., as described in PCT/US93/03868, hereby incorporated by reference in its entirety. Alternatively, e.g., when the prostate cancer sequence is up-regulated in prostate cancer, the activity of the endogenous prostate cancer gene is decreased, e.g., by the administration of a prostate cancer antisense nucleic acid.

In one embodiment, the prostate cancer proteins of the present invention may be used to generate polyclonal and monoclonal antibodies to prostate cancer proteins. Similarly, the prostate cancer proteins can be coupled, using standard technology, to affinity chromatography columns. These columns may then be used to purify prostate cancer antibodies useful for production, diagnostic, or therapeutic purposes. In a preferred embodiment, the antibodies are generated to epitopes unique to a prostate cancer protein; that is, the antibodies show little or no cross-reactivity to other proteins. The prostate cancer antibodies may be coupled to standard affinity chromatography columns and used to purify prostate cancer proteins. The antibodies may also be used as blocking polypeptides, as outlined above, since they will specifically bind to the prostate cancer protein.

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Methods of identifying variant prostate cancer-associated sequences

Without being bound by theory, expression of various prostate cancer sequences is correlated with prostate cancer or other prostate disorders. Accordingly, disorders based on mutant or variant prostate cancer genes may be determined. In one embodiment, the invention provides methods for identifying cells containing variant prostate cancer genes, e.g., determining all or part of the sequence of at least one endogenous prostate cancer genes in a cell. This may be accomplished using many sequencing techniques. In a preferred

embodiment, the invention provides methods of identifying the prostate cancer genotype of an individual, e.g., determining all or part of the sequence of at least one prostate cancer gene of the individual. This is generally done in at least one tissue of the individual, and may include the evaluation of a number of tissues or different samples of the same tissue. The method may include comparing the sequence of the sequenced prostate cancer gene to a known prostate cancer gene, e.g., a wild-type gene.

The sequence of all or part of the prostate cancer gene can then be compared to the sequence of a known prostate cancer gene to determine if differences exist. This can be done using many known homology programs, such as Bestfit, etc. In a preferred embodiment, the presence of a difference in the sequence between the prostate cancer gene of the patient and the known prostate cancer gene correlates with a disease state or a propensity for a disease state, as outlined herein.

In a preferred embodiment, the prostate cancer genes are used as probes to determine the number of copies of the prostate cancer gene in the genome.

In another preferred embodiment, the prostate cancer genes are used as probes to determine the chromosomal localization of the prostate cancer genes. Information such as chromosomal localization finds use in providing a diagnosis or prognosis in particular when chromosomal abnormalities such as translocations, and the like are identified in the prostate cancer gene locus.

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#### Administration of pharmaceutical and vaccine compositions

In one embodiment, a therapeutically effective dose of a prostate cancer protein or modulator thereof, is administered to a patient. By "therapeutically effective dose" herein is meant a dose that produces effects for which it is administered. The exact dose will depend on the purpose of the treatment, and will be ascertainable by one skilled in the art using known techniques (e.g., Ansel, et al. (1992) Pharmaceutical Dosage Forms and Drug Delivery; Lieberman (1993) Pharmaceutical Dosage Forms (vols. 1-3, Dekker, ISBN 0824770846, 082476918X, 0824712692, 0824716981; Lloyd (1999) The Art, Science and Technology of Pharmaceutical Compounding Amer. Pharma. Assn.; and Pickar (1999) Dosage Calculations Thomson). Adjustments for prostate cancer degradation, systemic versus localized delivery, and rate of new protease synthesis, as well as the age, body weight, general health, sex, diet, time of administration, drug interaction, and the severity of the

condition may be necessary, and will be ascertainable with routine experimentation by those skilled in the art. U.S. Patent Application N. 09/687,576 further discloses the use of compositions and methods of diagnosis and treatment in prostate cancer is hereby expressly incorporated by reference.

A "patient" for the purposes of the present invention includes both humans and other animals, particularly mammals. Thus the methods are applicable to both human therapy and veterinary applications. In the preferred embodiment the patient is a mammal, preferably a primate, and in the most preferred embodiment the patient is human. The patient typically will suffer from a prostate proliferative disorder, e.g., malignant or non-malignant, and may include cancer of other related conditions or disorders.

The administration of the prostate cancer proteins and modulators thereof of the present invention can be done in a variety of ways as discussed above, including, but not limited to, orally, subcutaneously, intravenously, intranasally, transdermally, intraperitoneally, intramuscularly, intrapulmonary, vaginally, rectally, or intraocularly. In some instances, e.g., in the treatment of wounds and inflammation, the prostate cancer proteins and modulators may be directly applied as a solution or spray, or via catheter.

The pharmaceutical compositions of the present invention comprise a prostate cancer protein in a form suitable for administration to a patient. In the preferred embodiment, the pharmaceutical compositions are in a water soluble form, such as being present as pharmaceutically acceptable salts, which is meant to include both acid and base addition salts. "Pharmaceutically acceptable acid addition salt" refers to those salts that retain the biological effectiveness of the free bases and that are not biologically or otherwise undesirable, formed with inorganic acids such as hydrochloric acid, hydrobromic acid, sulfuric acid, nitric acid, phosphoric acid and the like, and organic acids such as acetic acid, propionic acid, glycolic acid, pyruvic acid, oxalic acid, maleic acid, malonic acid, succinic acid, fumaric acid, tartaric acid, citric acid, benzoic acid, cinnamic acid, mandelic acid, methanesulfonic acid, ethanesulfonic acid, p-toluenesulfonic acid, salicylic acid, and the like. "Pharmaceutically acceptable base addition salts" include those derived from inorganic bases such as sodium, potassium, lithium, ammonium, calcium, magnesium, iron, zinc, copper, manganese, aluminum salts, and the like. Particularly preferred are the ammonium, potassium, sodium, calcium, and magnesium salts. Salts derived from pharmaceutically acceptable organic non-toxic bases include salts of primary, secondary, and tertiary amines,

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substituted amines including naturally occurring substituted amines, cyclic amines and basic ion exchange resins, such as isopropylamine, trimethylamine, diethylamine, triethylamine, tripropylamine, and ethanolamine.

The pharmaceutical compositions may also include one or more of the following: carrier proteins such as serum albumin; buffers; fillers such as microcrystalline cellulose, lactose, corn and other starches; binding agents; sweeteners and other flavoring agents; coloring agents; and polyethylene glycol.

The pharmaceutical compositions can be administered in a variety of unit dosage forms depending upon the method of administration. For example, unit dosage forms suitable for oral administration include, but are not limited to, powder, tablets, pills, capsules and lozenges. It is recognized that prostate cancer protein modulators (e.g., antibodies, antisense constructs, ribozymes, small organic molecules, etc.) when administered orally, should be protected from digestion. This is typically accomplished either by complexing the molecule(s) with a composition to render it resistant to acidic and enzymatic hydrolysis, or by packaging the molecule(s) in an appropriately resistant carrier, such as a liposome or a protection barrier. Means of protecting agents from digestion are well known in the art.

The compositions for administration will commonly comprise a prostate cancer protein modulator dissolved in a pharmaceutically acceptable carrier, preferably an aqueous carrier. A variety of aqueous carriers can be used, e.g., buffered saline and the like. These solutions are typically sterile and generally free of undesirable matter. These compositions may be sterilized by conventional, well known sterilization techniques. The compositions may contain pharmaceutically acceptable auxiliary substances as required to approximate physiological conditions such as pH adjusting and buffering agents, toxicity adjusting agents and the like, e.g., sodium acetate, sodium chloride, potassium chloride, calcium chloride, sodium lactate, and the like. The concentration of active agent in these formulations can vary widely, and will be selected primarily based on fluid volumes, viscosities, body weight, and the like in accordance with the particular mode of administration selected and the patient's needs (e.g., (1980) Remington's Pharmaceutical Science (15th ed.); and Hardman, et al. (eds. 2001) Goodman & Gilman: The Pharmacological Basis of Therapeutics McGraw-Hill.

Thus, a typical pharmaceutical composition for intravenous administration would be about 0.1 to 10 mg per patient per day. Dosages from 0.1 up to about 100 mg per patient per day may be used, particularly when the drug is administered to a secluded site and not into

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the blood stream, such as into a body cavity or into a lumen of an organ. Substantially higher dosages are possible in topical administration. Actual methods for preparing parenterally administrable compositions will be known or apparent to those skilled in the art, e.g., Remington's Pharmaceutical Science and Goodman and Gilman: The Pharmacological Basis of Therapeutics, supra.

The compositions containing modulators of prostate cancer proteins can be administered for therapeutic or prophylactic treatments. In therapeutic applications, compositions are administered to a patient suffering from a disease (e.g., a cancer) in an amount sufficient to cure or at least partially retard or arrest the disease and its complications. An amount adequate to accomplish this is defined as a "therapeutically effective dose." Amounts effective for this use will depend upon the severity of the disease and the general state of the patient's health. Single or multiple administrations of the compositions may be administered depending on the dosage and frequency as required and tolerated by the patient. The composition should provide a sufficient quantity of the agents of this invention to effectively treat the patient. An amount of modulator that is capable of preventing or slowing the development of cancer in a mammal is referred to as a "prophylactically effective dose." The particular dose required for a prophylactic treatment will depend upon the medical condition and history of the mammal, the particular cancer being prevented, as well as other factors such as age, weight, gender, administration route, efficiency, etc. Such prophylactic treatments may be used, e.g., in a mammal who has previously had cancer to prevent a recurrence of the cancer, or in a mammal who is suspected of having a significant likelihood of developing cancer, e.g., based partly on gene expression profiles.

It will be appreciated that the present prostate cancer protein-modulating compounds can be administered alone or in combination with additional prostate cancer modulating compounds or with other therapeutic agent, e.g., other anti-cancer agents or treatments.

In numerous embodiments, one or more nucleic acids, e.g., polynucleotides comprising nucleic acid sequences set forth in Tables 1A-4such as antisense polynucleotides, silencing RNA, or ribozymes, will be introduced into cells, in vitro or in vivo. The present invention provides methods, reagents, vectors, and cells useful for expression of prostate cancer-associated polypeptides and nucleic acids using in vitro (cell-free), ex vivo or in vivo (cell or organism-based) recombinant expression systems.

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The particular procedure used to introduce the nucleic acids into a host cell for expression of a protein or nucleic acid is application specific. Many procedures for introducing foreign nucleotide sequences into host cells may be used. These include the use of calcium phosphate transfection, spheroplasts, electroporation, liposomes, microinjection, plasma vectors, viral vectors, and many other well known methods for introducing cloned genomic DNA, cDNA, synthetic DNA, or other foreign genetic material into a host cell (see, e.g., Berger and Kimmel (1987) Guide to Molecular Cloning Techniques from Methods in Enzymology (vol. 152) Academic Press; Ausubel, et al., (eds. supplemented through 1999) Current Protocols Lippincott; and Sambrook, et al. (1989) Molecular Cloning: A Laboratory Manual (2d ed., Vol. 1-3) CSH Press.

In a preferred embodiment, prostate cancer proteins and modulators are administered as therapeutic agents, and can be formulated as outlined above. Similarly, prostate cancer genes (including both the full-length sequence, partial sequences, or regulatory sequences of the prostate cancer coding regions) can be administered in a gene therapy application. These prostate cancer genes can include antisense applications, either as gene therapy (i.e., for incorporation into the genome) or as antisense compositions, as will be appreciated by those in the art.

Prostate cancer polypeptides and polynucleotides can also be administered as vaccine compositions to stimulate HTL, CTL, and antibody responses.. Such vaccine compositions can include, e.g., lipidated peptides (see, e.g., Vitiello, et al. (1995) J. Clin. Invest. 95:341-349), peptide compositions encapsulated in poly(DL-lactide-co-glycolide) ("PLG") microspheres (see, e.g., Eldridge, et al. (1991) Molec. Immunol. 28:287-294; Alonso, et al. (1994) Vaccine 12:299-306; Jones, et al. (1995) Vaccine 13:675-681), peptide compositions contained in immune stimulating complexes (ISCOMS) (see, e.g., Takahashi, et al. (1990) Nature 344:873-875; Hu, et al. (1998) Clin Exp Immunol. 113:235-243), multiple antigen peptide systems (MAPs) (see, e.g., Tam (1988) Proc. Natl. Acad. Sci. USA 85:5409-5413; Tam (1996) J. Immunol. Methods 196:17-32), peptides formulated as multivalent peptides; peptides for use in ballistic delivery systems, typically crystallized peptides, viral delivery vectors (Perkus, et al., p. 379, in Kaufmann (ed. 1996) Concepts in vaccine development de Gruyter; Chakrabarti, et al. (1986) Nature 320:535-537; Hu, et al. (1986) Nature 320:537-540; Kieny, et al. (1986) AIDS Bio/Technology 4:790-xxx; Top, et al. (1971) J. Infect. Dis. 124:148-154; Chanda, et al. (1990) Virology 175:535-547), particles of viral or synthetic

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origin (see, e.g., Kofler, et al. (1996) J. Immunol. Methods 192:25-35; Eldridge, et al. (1993) Sem. Hematol. 30:16-24; Falo, et al. (1995) Nature Med. 7:649-653), adjuvants (Warren, et al. (1986) Annu. Rev. Immunol. 4:369-388; Gupta, et al. (1993) Vaccine 11:293-306), liposomes (Reddy, et al. (1992) J. Immunol. 148:1585-1589; Rock (1996) Immunol. Today 17:131-137), or, naked or particle absorbed cDNA (Ulmer, et al. (1993) Science 259:1745-1749; Robinson, et al. (1993) Vaccine 11:957-960; Shiver, et al., p. 423, in Kaufmann (ed. 1996) Concepts in Vaccine Development de Gruyter; Cease and Berzofsky (1994) Annu. Rev. Immunol. 12:923-989; and Eldridge, et al. (1993) Sem. Hematol. 30:16-24). Toxintargeted delivery technologies, also known as receptor mediated targeting, such as those of Avant Immunotherapeutics, Inc. (Needham, Massachusetts) may also be used.

Vaccine compositions often include adjuvants. Many adjuvants contain a substance designed to protect the antigen from rapid catabolism, such as aluminum hydroxide or mineral oil, and a stimulator of immune responses, such as lipid A, Bortadella pertussis or Mycobacterium tuberculosis derived proteins. Certain adjuvants are commercially available as, e.g., Freund's Incomplete Adjuvant and Complete Adjuvant (Difco Laboratories, Detroit, MI); Merck Adjuvant 65 (Merck and Company, Inc., Rahway, NJ); AS-2 (SmithKline Beecham, Philadelphia, PA); aluminum salts such as aluminum hydroxide gel (alum) or aluminum phosphate; salts of calcium, iron or zinc; an insoluble suspension of acylated tyrosine; acylated sugars; cationically or anionically derivatized polysaccharides; polyphosphazenes; biodegradable microspheres; monophosphoryl lipid A, and quil A. Cytokines, such as GM-CSF, interleukin-2, -7, -12, and other like growth factors, may also be used as adjuvants.

Vaccines can be administered as nucleic acid compositions wherein DNA or RNA encoding one or more of the polypeptides, or a fragment thereof, is administered to a patient. This approach is described, for instance, in Wolff, et al. (1990) <u>Science</u> 247:1465-1468 as well as U.S. Patent Nos. 5,580,859; 5,589,466; 5,804,566; 5,739,118; 5,736,524; 5,679,647; WO 98/04720; and in more detail below. Examples of DNA-based delivery technologies include "naked DNA", facilitated (bupivicaine, polymers, peptide-mediated) delivery, cationic lipid complexes, and particle-mediated ("gene gun") or pressure-mediated delivery (see, e.g., U.S. Patent No. 5,922,687).

For therapeutic or prophylactic immunization purposes, the peptides of the invention can be expressed by viral or bacterial vectors. Examples of expression vectors include

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attenuated viral hosts, such as vaccinia or fowlpox. This approach involves the use of vaccinia virus, e.g., as a vector to express nucleotide sequences that encode prostate cancer polypeptides or polypeptide fragments. Upon introduction into a host, the recombinant vaccinia virus expresses the immunogenic peptide, and thereby elicits an immune response. Vaccinia vectors and methods useful in immunization protocols are described in, e.g., U.S. Patent No. 4,722,848. Another vector is BCG (Bacille Calmette Guerin). BCG vectors are described in Stover, et al. (1991) Nature 351:456-460. A wide variety of other vectors useful for therapeutic administration or immunization, e.g., adeno and adeno-associated virus vectors, retroviral vectors, Salmonella typhi vectors, detoxified anthrax toxin vectors, and the like, will be apparent to those skilled in the art from the description herein (see, e.g., Shata, et al. (2000) Mol. Med. Today 6:66-71; Shedlock, et al. (2000) J. Leuk. Biol. 68:793-806; Hipp, et al. (2000) In Vivo 14:571-85).

Methods for the use of genes as DNA vaccines are well known, and include placing a prostate cancer gene or portion of a prostate cancer gene under the control of a regulatable promoter or a tissue-specific promoter for expression in a prostate cancer patient. The prostate cancer gene used for DNA vaccines can encode full-length prostate cancer proteins, but more preferably encodes portions of the prostate cancer proteins including peptides derived from the prostate cancer protein. In one embodiment, a patient is immunized with a DNA vaccine comprising a plurality of nucleotide sequences derived from a prostate cancer gene. For example, prostate cancer-associated genes or sequence encoding subfragments of a prostate cancer protein are introduced into expression vectors and tested for their immunogenicity in the context of Class I MHC and an ability to generate cytotoxic T cell responses. This procedure may provide for production of cytotoxic T lymphocyte responses against cells which present antigen, including intracellular epitopes.

In a preferred embodiment, the DNA vaccines include a gene encoding an adjuvant molecule with the DNA vaccine. Such adjuvant molecules include cytokines that increase the immunogenic response to the prostate cancer polypeptide encoded by the DNA vaccine. Additional or alternative adjuvants are available.

In another preferred embodiment prostate cancer genes find use in generating animal models of prostate cancer. When the prostate cancer gene identified is repressed or diminished in cancer tissue, gene therapy technology, e.g., wherein antisense RNA directed to the prostate cancer gene will also diminish or repress expression of the gene. Animal

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models of prostate cancer find use in screening for modulators of a prostate cancer-associated sequence or modulators of prostate cancer. Similarly, transgenic animal technology including gene knockout technology, e.g., as a result of homologous recombination with an appropriate gene targeting vector, will result in the absence or increased expression of the prostate cancer protein. When desired, tissue-specific expression or knockout of the prostate cancer protein may be necessary.

It is also possible that the prostate cancer protein is overexpressed in prostate cancer. As such, transgenic animals can be generated that overexpress the prostate cancer protein. Depending on the desired expression level, promoters of various strengths can be employed to express the transgene. Also, the number of copies of the integrated transgene can be determined and compared for a determination of the expression level of the transgene. Animals generated by such methods find use as animal models of prostate cancer and are additionally useful in screening for modulators to treat prostate cancer.

#### 15 Kits for Use in Diagnostic and/or Prognostic Applications

For use in diagnostic, research, and therapeutic applications suggested above, kits are also provided by the invention. In the diagnostic and research applications such kits may include one of the following: assay reagents, buffers, prostate cancer-specific nucleic acids or antibodies, hybridization probes and/or primers, antisense polynucleotides, silencing RNA, ribozymes, dominant negative prostate cancer polypeptides or polynucleotides, small molecules inhibitors of prostate cancer-associated sequences, etc. A therapeutic product may include sterile saline or another pharmaceutically acceptable emulsion and suspension base.

In addition, the kits may include instructional materials containing instructions (i.e., protocols) for the practice of the methods of this invention. While the instructional materials typically comprise written or printed materials they are not limited to such. A medium capable of storing such instructions and communicating them to an end user is contemplated by this invention. Such media include, but are not limited to electronic storage media (e.g., magnetic discs, tapes, cartridges, chips), optical media (e.g., CD ROM), and the like. Such media may include addresses to internet sites that provide such instructional materials.

The present invention also provides for kits for screening for modulators of prostate cancer-associated sequences. Such kits can be prepared from readily available materials and reagents. For example, such kits can comprise one or more of the following materials: a

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prostate cancer-associated polypeptide or polynucleotide, reaction tubes, and instructions for testing prostate cancer-associated activity. Optionally, the kit contains biologically active prostate cancer protein. A wide variety of kits and components can be prepared according to the present invention, depending upon the intended user of the kit and the particular needs of the user. Diagnosis would typically involve evaluation of a plurality of genes or products. The genes will be selected based on correlations with important parameters in disease which may be identified in historical or outcome data.

# **EXAMPLES**

Example 1: Gene Chip Analyses of Expression Profiles

Molecular profiles of various normal and cancerous tissues were determined and analyzed using gene chips. RNA was isolated and gene chip analysis was performed as described (Glynne, et al. (2000) Nature 403:672-676; Zhao, et al. (2000) Genes Dev. 14:981-993).

### EXAMPLE 2: Identification of androgen dependent/independent genes

To identify gene expression changes during the transition from androgen-dependent to androgen-independent prostate cancer, oligonucleotide microarrays ("K" chips or Affymetrix Eos Hu03) were interrogated with cRNAs derived from the human CWR22 prostate cancer xenograft model propagated in nude mice (Pretlow, et al. (1993) J. Natl. Cancer Inst. 85:394-398). The CWR22 xenograft is androgen-dependent when grown in male Nude mice. Androgen-independent sub-lines can be derived by first establishing androgen-dependent tumors in male mice. The mice are then castrated to remove the primary source of growth stimulus (androgen), resulting in tumor regression. Within 3-10 months molecular events prompt the tumors to relapse and start growing as androgen-independent tumors. See, e.g., Nagabhushan, et al. (1996) Cancer Res. 56:3042-3046; Amler, et al. (2000) Cancer Res. 60:6134-6141; and Bubendorf, et al. (1999) J. Natl. Cancer Inst. 91:1758-1764.

Using the CWR22 xenograft model, tumors were grown subcutaneously in male nude mice. Tumors were harvested at different times after castration. The time points post-castration included (in days): 0, 1, 3, 4, 5, 10, 30, 40, 50, 51, 52, 59, 60, 61, 70, 79, 80, 82, 120, and 125. Analyses also included established androgen-independent xenografts. Castration resulted in tumor regression. At day 120 and thereafter, the tumors relapsed and started growing in the absence of androgen.

cRNAs were generated by in vitro transcription assays (IVTs) from the different samples and were hybridized to the oligonucleotide microarrays (Affymetrix Eos Hu03). Hybridization was measured by the average fluorescence intensity (AI), which is directly proportional to the expression level of the gene.

Two types of analyses were applied to the results:

Analysis A:

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The samples were divided into different time groups which included the following time points post castration (in days): 1-5, 10, 30-40, 50-82, 120-125. To identify changes in gene expression, the following calculations were made:

- 1. The median (or mean, in case there were only 2 samples in a group) was calculated for each group.
  - 2. The medians (or means) for each group was compared to one-another.
  - 3. Genes were selected that exhibited a minimum 2 fold difference in the median (or mean) between any of the groups.
- 4. The change in gene expression over time was analyzed for each selected gene to look for specific pattern changes.

Only genes with an interesting expression pattern during the androgen-ablation time course were selected as potential new therapeutic targets and/or diagnostic markers. Among the 70,000 gene clusters present on Hu01 and Hu02, we identified 820 gene clusters with the desired expression patterns. These expression patterns can be broadly defined into the following categories:

- 1. Genes that are expressed early in the time course, then drop off in expression, and then express again with emergence of androgen-independence (hi-lo-hi pattern in Table 1A).
- 2. Genes that are expressed early in the time course, then drop off in expression, and do not express again with emergence of androgen-independence (hi-lo-lo pattern in Table 1A).
- 20 3. Genes that are not expressed early in the time course, but express only with emergence of androgen-independence (lo-lo-hi pattern in Table 1A).
  - 4. Genes that are not expressed early in the time course, but then express as androgen is withdrawn and continue to express with emergence of androgen-independence (lo-hi-hi pattern in Table 1A).
- 25 5. Genes that are not expressed early in the time course, but then express as androgen is withdrawn and drop off again with emergence of androgen-independence (lo-hi-lo pattern in Table 1A).

Group 1 is characterized by cell-cycle regulating genes, such as those encoding cyclin B1, p21/WAF1, CDC18-homolog, cyclin A2, cyclin D1, and possible growth factors such as hAG2 (anterior gradient 2 homolog) among others. This indicates that interruption of growth factor and/or cell cycle pathways prevents the emergence of androgen-independent disease, making group 1 genes good targets for treating advanced prostate cancer.

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Group 2 represents genes that are androgen-dependent, and do not re-express due to the lack of androgen signal in the androgen-independent phenotype. This group includes genes encoding proteins such as Fibronectin 1, which has been previously shown to be down-regulated with androgen-withdrawal (Amler, et al. (2000) <u>Cancer Res.</u> 60:6134-6141).

Group 3 represents genes that are up-regulated by signals that induce the androgen-independent phenotype. This group includes genes encoding stanniocalcin 2, c-fos proto-oncogene product, vascular endothelial growth factor, the cell surface protein transmembrane 4 superfamily member 1 and adrenomedullin among others. Adrenomedullin has recently been shown to act as an autocrine growth factor for the androgen-independent prostate cancer cell line DU145 (Rocchi, et al. (2001) Cancer Res. 61:1196-1206), indicating that its up-regulation is critical for supporting an androgen-independent phenotype. Blocking adrenomedullin function, and/or other genes in this group, prevents the growth of androgen-independent tumor cells.

Group 4 represents genes that are androgen-repressed and are only expressed in the absence of androgen. This group includes genes encoding the protein tyrosine phosphatase interacting protein liprin-alpha 2, the CD24 antigen, and the catalytic subunit for phosphatidylinositol 4-kinase amongst others. Patients that are treated for advanced prostate cancer by hormone-ablation may have in their bodies cells that have survived hormone-ablation and are likely to up-regulate genes that belong to Group 4. Therefore, Group 4 gene products are particularly good therapeutic targets for treating patients undergoing hormone-ablation therapy.

Group 5 represents genes that are involved in regulating signals that induce an androgen-independent phenotype. This group includes genes encoding Rab2 (a Ras-like G protein), the Son of Sevenless homolog (a GTP/GDP exchange factor involved in activating Ras-like proteins), and the p85 regulatory subunit for phosphoinositide-3-kinase (PI3-kinase). The PI3-kinase pathway has been implicated in providing a survival signal to the prostate cancer cell line LNCaP (Lin, et al. (1999) Cancer Res. 59:2891-2897). This indicates that ras-like signals and signals dependent on PI3-kinase are involved in inducing the androgen-independent phenotype. For that reason, Group 5 gene products are particularly good therapeutic targets for treating patients undergoing hormone-ablation therapy. Analysis B:

For the second analysis, the samples were divided into 4 time groups which included the following time points post castration (in days): 0-1, 3-5, 10-82, >120. To identify changes in gene expression, the following analysis was performed:

- 1. Genes were selected that exhibited a minimum of 100 AI units at the 90<sup>th</sup> percentile expression level of samples.
- 2. The group mean expression levels for each gene were calculated. The genes were further sub-selected to exhibit a minimum 3 fold difference between the group means.
- 3. An analysis of variance was then performed on selected genes. From the original 59,680 gene clusters present on the Hu03 gene chip, only about 1165 genes with a P value of < 0.01 were identified that also exhibited the above mentioned parameters.
- 4. A method was then employed for calculating the positive false discovery rate (pFDR), i.e., an estimate of the proportion of false-positives present in a set of findings (Storey and Tibshirani (2001) Technical Report, Department of Statistics, Stanford University, CA). This technique was developed explicitly for use with microarray data. The procedure involves randomly assigning the membership status of each sample to a group and reperforming the analysis of variance. In each simulation, the number of group members (6 for Group 1, 9 for group 2, 15 for group 3, and 4 for group 4) remained constant, but these designations were shuffled and assigned to each sample at random. The permutation was

performed 1000 times, and for each simulation, the number of findings at P < 0.01 was noted.

- The number of false positives under null conditions, was then divided by the number of actual findings (n=1165 genes) to obtain an estimate of the proportion of false positive findings. After the application of a correction factor, the final estimate for the pFDR was about 1%. Thus, one can expect that approximately 12 of the 1165 findings are false positives.
- 5. The approximately 1165 genes were clustered by expression pattern to identify specific pattern changes. Only genes with an interesting expression pattern during the androgenablation time course were selected as potential new therapeutic targets and/or diagnostic markers. These expression patterns can be broadly defined into the following categories:
  - 1. Genes that are expressed early in the time course of androgen withdrawal, then drop off in expression, and then express again with emergence of androgen-independence (hi-lo-lo-hi pattern in Table 2A).

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2. Genes that are expressed early in the time course, then drop off in expression immediately after androgen-withdrawal, and do not express again with emergence of androgen-independence (hi-lo-lo-lo pattern in Table 2A).

- 3. Genes that are expressed early in the time course, then drop off in expression after several days of androgen withdrawal, and do not express again with emergence of androgen-independence (hi-hi-lo-lo pattern in Table 2A).
- 4. Genes that are not expressed early in the time course, but express only with emergence of androgen-independence (lo-lo-lo-hi pattern in Table 2A).
- 5. Genes that are not expressed early in the time course, but then express as androgen is withdrawn and continue to express with emergence of androgen-independence (lo-lo-hi-hi pattern in Table 2A).
  - 6. Genes that are not expressed early in the time course, but then express as androgen is withdrawn and drop off again with emergence of androgen-independence (lo-lo-hi-lo pattern in Table 2A).

Group 1 is characterized by cell-cycle regulating genes and cell growth promoting genes, such as those encoding cyclin B1 and CDC45 among others, growth factors/hormones such as hAG2 (anterior gradient 2 homolog), adrenomedullin, and stanniocalcin 2 among others, and growth factor receptors, such as the bone morphogenic protein receptor type 1B (BMP-R1B) and the endothelial differentiation lysophosphatidic acid G-protein-coupled receptor 7 among others. Adrenomedullin has recently been shown to act as an autocrine growth factor for the androgen-independent prostate cancer cell line DU145 (Rocchi, et al. (2001) Cancer Res. 61:1196-1206), indicating that its up-regulation is critical for supporting an androgen-independent phenotype. This indicates that interruption of growth factor and/or cell cycle pathways prevents the emergence of androgen-independent disease, making group 1 genes good targets for treating both localized and advanced prostate cancer and related conditions.

Group 2 represents genes that are androgen-dependent, and do not re-express due to the lack of androgen signal in the androgen-independent phenotype. This group includes genes encoding proteins such as the endothelial protein C receptor (EPCR) and the potassium intermediate/small conductance calcium-activated channel (subfamily N, member 2). These genes represent targets for treating androgen-dependent prostate cancer and related conditions.

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Group 3 also represents genes that are androgen-dependent, and do not re-express due to the lack of androgen signal in the androgen-independent phenotype. This group includes genes encoding proteins such as Fibronectin 1, which has been previously shown to be down-regulated with androgen-withdrawal (Amler, et al. (2000) <u>Cancer Res.</u> 60:6134-6141), and genes encoding signaling proteins such as Rho GTPase activating protein 1. These genes represent targets for treating androgen-dependent prostate cancer and related conditions.

Group 4 represents genes that are up-regulated by signals that induce and maintain the androgen-independent phenotype. This group includes genes encoding potential growth promoting proteins such as chemokine-like factor (Unigene ID Hs.15159), colon cancerassociated protein Mic1, and the mitogen-activated protein kinase-activated protein kinase 2. Blocking function of these proteins, and/or other genes in this group, prevents the growth of androgen-independent tumor cells and related conditions.

Group 5 represents genes that are androgen-repressed and are only expressed in the absence of androgen or that are induced by the absence of androgen. This group includes genes encoding transcriptional regulators such as the androgen receptor, the DNA activated protein kinase (catalytic subunit), and nuclear factor related to kappa B binding protein (NFRKB), among others. Patients that are treated for advanced prostate cancer by hormone-ablation may have in their bodies cells that have survived hormone-ablation and are likely to up-regulate genes that belong to Group 5. Therefore, Group 5 gene products are particularly good therapeutic targets for treating patients undergoing hormone-ablation therapy.

Group 6 represents genes that are involved in regulating signals that are induced during androgen withdrawal and that induce an androgen-independent phenotype. This group includes genes encoding signaling molecules such as phosphoinositide-3-kinase (class 2, alpha polypeptide), signal transducer and activator of transcription 2 (STAT2), phospholipase A2 (group IIA) and the protein tyrosine phosphatase interacting protein liprin-alpha 2, cell surface receptors such as gamma-aminobutyric acid (GABA) A receptor epsilon subunit, G protein-coupled receptor 48, and immune function proteins such as the major histocompatibility complex class II DR alpha. The PI3-kinase pathway has been implicated in providing a survival signal to the prostate cancer cell line LNCaP (Lin, et al. (1999) Cancer Res. 59:2891-2897). This indicates that ras-like signals and signals dependent on PI3-kinase are involved in inducing the androgen-independent phenotype. For that reason, Group 6 gene

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products are particularly good therapeutic targets for treating patients undergoing hormone-ablation therapy.

TABLE 1A provides Accession numbers for genes, including expressed sequence tags, (incorporated in their entirety here and throughout the application where Accession numbers are provided). Genes with an interesting expression pattern during the androgen-ablation time course were selected as potential new therapeutic targets and/or diagnostic markers. 820 gene clusters were identified with desired expression patterns. These expression patterns can be broadly defined into the following categories:

1. Genes that are expressed early in the time course, then drop off in expression, and then express again with emergence of androgen-independence (hi-lo-lo pattern).

2. Genes that are expressed early in the time course, but express only with emergence of androgen-independence (lo-lo-hi pattern).

3. Genes that are not expressed early in the time course, but express only with emergence of androgen-independence (lo-lo-hi pattern).

4. Genes that are not expressed early in the time course, but then express as androgen is withdrawn and continue to express with emergence of androgen-independence (lo-hi-hi nattern).

- 5. Genes that are not expressed early in the time course, but then express as androgen is withdrawn and drop off again with emergence of androgen-independence (lo-hi-lo 10 pattern).

Table 1B lists accession numbers for primekeys lacking a unigenelD in table 1A. For each probeset is listed a gene cluster number from which oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

Table 1C lists genomic positioning for primekeys lacking unigene ID's and accession numbers in tables 1A. For each predicted exon is listed genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed.

	TABLE 44				
20	TABLE 1A Pkey	ExAccn	UnigenelD	Unigene Title	pattern
20	,		Unigenero	oragene rad	pattern
	102772	U83115	Hs.161002	absent in metanoma 1	hi-lo-hi
	128610	N48373	Hs.10247	activated leucocyte cell adhesion molecu	hi-lo-hi
0.5	102276	N48373	Hs.10247	activated leucocyte cell adhesion molecu	hi-lo-hi
25	100654	A03758			hi-lo-hi
	100655	A03758			hi-lo-hi
	135400	X78592	Hs.99915	androgen receptor (dihydrotestosterone r	hi-lo-hi
	331363	AW582256	"Hs.91011	anterior gradient 2 (Xenepus laevis) hom	hi-to-hi
20	115764	AW582256	'Hs.91011	anterior gradient 2 (Xenepus taevis) hom	hi-to-hi
30	120483	BE251623	Hs.1578	baculoviral IAP repeat-containing 5 (sur	hi-lo-hi
	101505	AA307680	Hs.75692	asparagine synthetase	hi-lo-hi
	127236	AW661857	Hs.98658	budding uninhibited by benzimidazoles 1	hì-lo-hi
	128472	BE241880	"Hs.10029	cathepsin C	hi-lo-hi
35	102712	U77949	Hs.69563	CDC6 (cell division cycle 6, S. cerevisi	hì-lo-hi
55	314943 102123	Y00272	Hs.184572 "Hs.1594	cell division cycle 2, G1 to S and G2 to	hi-lo-hi
	326213	NM_001809	ns. 1594	centromere protein A (17kD)	hi-lo-hi
	327110			CH.17_hs gi[5867224 CH.21_hs gi[6117842	hi-lo-hi hi-lo-hi
	339186			CH22_DA59H18.GENSCAN,72-13	hi-lo-hi
40	337755			CH22_EM:AC000097.GENSCAN.109-2	hi-lo-hi
	337674			CH22_EM:AC000097.GENSCAN.67-4	hi-io-hi
	337675			CH22_EM:AC000097.GENSCAN.67-6	hi-lo-hi
	333516			CH22_FGENES.173_1	hi-lo-hi
	333517			CH22_FGENES.173_2	hi-lo-hi
45	333795			CH22_FGENES.275_1	hi-lo-hi
	333796			CH22_FGENES.275_3	hi-lo-hi
	333808			CH22_FGENES.279_2	hi-lo-hi
	333809			CH22_FGENES.280_2	hi-lo-hi
50	-332792			CH22_FGENES.3_2	hi-lo-hi
50	334101			CH22_FGENES.327_59	hi-lo-hi
•	334502			CH22_FGENES.397_18	hi-lo-hi
	334616			CH22_FGENES.411_15	hi-lo-hi
	334899			CH22_FGENES.452_13	hi-lo-hi
55	334900			CH22_FGENES.452_14	hi-lo-hi
55	334902			CH22_FGENES.452_16	hi-lo-hi
	334905 334906			CH22_FGENES.452_20	hi-lo-hi hi-lo-hi
	334951			CH22_FGENES.452_21 CH22_FGENES.465_20	hi-lo-hi
	335044			CH22_FGENES.480_1	hi-lo-hi
60	335753			CH22_FGENES.604_2	hi-lo-hi
	335755			CH22_FGENES.604_4	hi-lo-hi
	333135			CH22_FGENES.83_11	hì-lo-hi
	333137			CH22_FGENES.83_13	hi-lo-hi
	333138			CH22_FGENES.83_15	hi-lo-hi
65	333139			CH22_FGENES.83_16	hi-lo-hi
	336721			CH22_FGENES.83-17	hi-lo-hi
	105012	AF098158	Hs.9329	chromosome 20 open reading frame 1	hì-lo-hi
	134470	X54942	Hs.83758	CDC28 protein kinase 2	hi-lo-hi
70	134750	L29073	Hs.1139	cold shock domain protein A	hi-lo-hi
70	125819	AA044840	"Hs.251871	CTP synthase	hi-lo-hi
	102993	BE262998	Hs.85137	cyclin A2	hi-lo-hi
	131185	BE280074	Hs.23960	cyclin B1	hi-lo-hi
	106350 103080	AK001404 AU077231	"Hs.194698 "Hs.82932	cyclin B2 cyclin D1 (PRAD1: parathyroid adenomatos	hi-lo-hi
75	103080	AA284166	Hs.84113	cyclin-dependent kinase inhibitor 3 (CDK	hi-lo-hi hi-lo-hi
, 5	100589	AW247430	Hs.84152	cystathionine-beta-synthase	ni-to-ni hi-lo-hi
	130655	AI831962	Hs.17409	cysteine-rich protein 1 (intestinal)	ni-lo-ni hi-lo-hi
	101473	M22976	Hs.83834	cytochrome b-5	ni-to-ti hi-lo-hi
	101468	BE538296	"Hs.181028	cytochrome c oxidase subunit Va	hi-lo-hi
80	103546	Z14244	"Hs.75752	cytochrome c oxidase subunit VIIb	hi-lo-hi
	100829	AA471098	Hs.278544	acetyl-Coenzyme A acetyltransferase 2 (a	hi-lo-hi
	102469	AF058293	Hs.180015	O-dopachrome tautomerase	hi-ko-hi

5

	44.4000				
	114292	AI815395	Hs.184641	fatty acid desaturase 2	hi-lo-hi
	100656	BE250162	"Hs.83765	dihydrofolate reductase	hi-lo-hi
	133799	W24087	Hs.76285	DKFZP564B167 protein	hi-lo-hi
5	129113	BE543205	*Hs.288771	DKFZP586A0522 protein	hi-lo-hi
)	332732	AF191019	Hs.8361	hypothetical protein, estradiol-induced	hi-lo-hi
	108846	AL117452	*Hs.44155	DKFZP586G1517 protein	hi-lo-hi
	133903	X63692	"Hs.77462	DNA (cytosine-5-)-methyltransferase 1	hi-lo-hi
	320099	AW411307	Hs.114311	CDC45 (cell division cycle 45, S.cerevis	hi-lo-hi
10	321960	AA723883	Hs.302446	hypothetical protein MGC10334	hi-lo-hi
10	324988	AK001379	"Hs.121028	hypothetical protein FLJ10549	hi-lo-hi
	303274	AK001468	Hs.62180	anillin (Drosophila Scraps homolog), act	hi-lo-hi
	301804	AK001468	Hs.62180	anillin (Drosophila Scraps homolog), act	hi-lo-hi
	300551 304541	AW408800	Hs.104859 Hs.169476	hypothetical protein DKFZp762E1312	hi-lo-hi
15		AA482561	HS.109470	glyceraldehyde-3-phosphate dehydrogenase	hi-lo-hi
13	304521	AA464716	NI- 00700	gb:zx82c11.s1 Soares ovary tumor NbHOT H	hi-lo-hi
	129075	BE250162	"Hs.83765	dihydrofolate reductase	hi-lo-hi
	111003	N52980	Hs.83765	dihydrofolate reductase	hi-lo-hi
	115536	AK001468	Hs.62180	aniilin (Drosophila Scraps homolog), act	hi-lo-hi
20	108857	AK001468	Hs.62180	anillin (Drosophila Scraps homolog), act	hi-lo-hi
20	332397	AB027249	Hs.104741	PDZ-binding kinase; T-cell originated pr	hi-lo-hi
	330714	AA263143	Hs.24596	RAD51-interacting protein	hi-lo-hi
	104636	R82252	Hs.106106 Hs.22971	Homo sapiens cAMP-dependent protein kina ESTs	hi-lo-hi
	104986	AW088826		ESTs .	hi-lo-hi
25	105076	AI598252	Hs.37810		hi-lo-hi
23	105312	BE613348	"Hs.23348	S-phase kinase-associated protein 2 (p45	hì-lo-hi
	105388	AW575008	Hs.11355	thymopoietin	hi-lo-hi
	105953	BE410556	Hs.236556	hypothetical protein STRAIT11499	hi-lo-hi
	106286	AI765107	"Hs.274422	hypothetical protein FLJ20550	hi-lo-hi
30	106889	U46258	Hs.18349	HSPC145 protein	hi-lo-hi
50	109220 113158	AW958181	Hs.189998	ESTs	hi-lo-hi
	114542	AA328102 AW970128	Hs.24641 "Hs.293380	cytoskeleton associated protein 2 ESTs	hi-lo-hi
	114986	AK000361	Hs.133260	hypothetical protein FLJ20354	hi-lo-hi hi-lo-hi
	115291	BE545072	"Hs.122579	hypothetical protein FLJ10461	hi-lo-hi
35	115414	AA662240	Hs.283099	AF15q14 protein	hi-lo-hi
55	115471	AK001376	Hs.59346	hypothetical protein FLJ10514	hi-lo-hi
	115522	BE614387	Hs.47378	ESTs, Moderately similar to T50635 hypot	hi-lo-hi
	115652	BE093589	Hs.38178	hypothetical protein FLJ23468	hi-lo-hi
	116121	AK001330	Hs.48855	hypothetical protein FLJ 10468	hi-lo-hi
40	116130	AW183533	Hs.38178	hypothetical protein FLJ23468	hì-lo-hi
-10	116448	BE268321	Hs.208912	hypothetical protein MGC861	hi-lo-hi
	116787	AW362955	Hs.15641	ESTs	hi-lo-hi
	118336	BE327311	Hs.47166	HT021	hi-lo-hi
	120649	AA687322	Hs.192843	leucine zipper protein FKSG14	hi-lo-hi
45	121503	AA412049	Hs.290347	ESTs	hi-lo-hi
	121748	BE536911	Hs.234545	Homo sapiens NUF2R mRNA, complete cds	hi-lo-hi
,	122860	AA464414		gb:zx78g01.s1 Soares ovary tumor NbHOT H	hi-lo-hi
	123477	AF217515	Hs.283532	uncharacterized bone marrow protein BM03	hi-lo-hi
	130338	AJ375726	"Hs.279918	hypothetical protein	hi-lo-hi
50	130680	BE567313	Hs.183109	monoamine oxidase A	hi-lo-hi
	131148	AW953575	"Hs.303125	p53-induced protein PIGPC1	hi-lo-hi
	131626	BE514605	"Hs.289092	Homo sapiens cDNA: FLJ22380 fis, clone H	hi-lo-hi
	131937	AI907735	Hs.21446	Homo sapiens mRNA for KIAA1716 protein,	hi-lo-hi
	131965	W79283	Hs.35962	ESTs	hi-lo-hi
55	132371	AA235448	Hs.46677	PRO2000 protein	hi-lo-hi
	133626	AW836130	Hs.75277	hypothetical protein FLJ13910	hi-lo-hi
	300942	AW301344	Hs.122908	Homo sapiens, clone IMAGE:3048353, mRNA,	hi-lo-hi
	300953	AA542845	Hs.294088	ESTs	hi-lo-hi
60	302656	BE090580	Hs.70704	Homo sapiens, clone IMAGE:2823731, mRNA,	hi-lo-hi
60	311928	T62216	Hs.270840	ESTs	hi-lo-hi
	313637	AK000742	Hs.126774	L2DTL protein	hi-lo-hi
	313832	AW271106	Hs.133294	ESTs	hi-lo-hi
	316465	AW574774	Hs.121692	ESTs	hi-to-hi
c =	317202	AA894880	Hs.181181	ESTs	hi-lo-hi
65	320771	R74441	Hs.117176	poly(A)-binding protein, nuclear 1	hi-lo-hi
	321636	Al820961	Hs.193465	ESTs	hi-lo-hi
	330867	AW978991	Hs.221197	ESTs	hi-lo-hi
	331442	H77381	Hs.159420	ESTs	hi-lo-hi
70	106654	AW075485	Hs.286049	phosphoserine aminotransferase	hi-lo-hi
70	106590	Al350260	Hs.301539	hypothetical protein MGC2633	hi-lo-hi
	128460	T16206	Hs.237164	ESTs, Highly similar to LDHH_HUMAN L-LA	hi-lo-hi
	114394	T34462	Hs.103291	neuritin	hi-lo-hi
	315936	AW069807	Hs.271252	ESTs	hi-lo-hi
75	108886	AW248434	Hs.91521	hypothetical protein	hi-lo-hi
75	129241	A1878857	Hs.109706	hematological and neurological expressed	hi-lo-hi
	104978	Al199268	Hs.19322	ESTs, Wealdy similar to CGHU7L collagen	hi-lo-hi
	129626	F13272	Hs.111334	ferritin, light polypeptide	hi-lo-hi
	118895	BE304917	Hs.31097	hypothetical protein FLJ21478	hi-lo-hi
80	332577	Al826268	Hs.27769	ESTs, Weakly similar to MCAT_HUMAN MITOC	hi-lo-hi
90	116732	AW152225 Al216748	Hs.165909	ESTs ESTs, Wealdy similar to AF151859 1 CGI-1	hi-lo-hi
	106774	Al216748 BE612676	Hs.14587 Hs.303116	stromal cell-derived factor 2-like 1	hi-lo-hi bi lo bi
	108818	00012010	113.303110	Submidit Celeatived iopidit Kalife I	hi-lo-hi

	045040	41007044	FU- 454000	h17711 feet 11 4	L! I= L!
	315618	Al287341	"Hs.154029	bHLH factor Hes4	hi-lo-hi
	110561	AA379597	Hs.5199	HSPC150 protein similar to ubiquitin-con	hi-lo-hi
	132959	AW014195	Hs.61472	ESTs, Weakly similar to unknown [S.cerev	hi-lo-hi
_	103195	AA351647	Hs.2642	eukaryotic translation elongation factor	hi-lo-hi
5	100368	D79987	Hs.153479	extra spindle poles, S. cerevisiae, homo	hi-lo-hi
	103177	BE244377	*Hs.48876	famesyl-diphosphate famesyltransferase	hi-lo-hi
	109141	AF174600	Hs.193380	F-box protein Fbx20	hi-lo-hi
	100676	X02761	"Hs.287820	fibronectin 1	hi-lo-hi
10	100254	AA452181	Hs.77643	FK506-binding protein 1B (12.6 kD)	hi-lo-hi
10	133688	U71321	Hs.7557	FK506-binding protein 5	hi-lo-hi
	107129	AC004770	"Hs.4756	flap structure-specific endonuclease 1	hi-lo-hi
	102696	BE540274	Hs.239	forkhead box M1	hi-lo-hi
	101753	L11144	Hs.1907	galanin	hi-lo-hi
	101597	AA317089	"Hs.597	glutamic-oxaloacelic transaminase 1, sol	hi-lo-hi
15	133512	L18861		gb:Human Golli-mbp gene, exon 1.	hi-lo-hi
	130080	X14850	Hs.147097	H2A histone family, member X	hi-lo-hi
	101600	BE561617	"Hs.119192	H2A histone family, member Z	hi-lo-hi
	101332	J04088	"Hs.156346	topoisomerase (DNA) II alpha (170kD)	hi-lo-hi
	132967	AA316181	Hs.61635	six transmembrane epithelial antigen of	hì-lo-hi
20	129726	H15474	Hs.132898	falty acid desaturase 1	hi-lo-hi
20	106925	AK002011	Hs.37558		hi-lo-hi
				hypothetical protein FLJ11149	hi-lo-hi
	105643	BE621719	Hs.173802	KIAA0603 gene product	
	116028	H59799	Hs.42644	thioredoxin-like	hi-lo-hi
25	105437	AF151076	Hs.25199	hypothetical protein	hi-lo-hi
25	122512	AF053305	Hs.98658	budding uninhibited by benzimidazoles 1	hi-lo-hi
	131991	AF053306	Hs.36708	budding uninhibited by benzimidazoles 1	hi-lo-hi
	135015	AW361638	Hs.278338	LGN protein	hi-lo-hi
	102208	U22961		gb:Human mRNA clone with similarity to L	hi-lo-hi
	100144	AL119964	Hs.75616	seladin-1	hi-lo-hi
30	100447	NM_014767	Hs.74583	KIAA0275 gene product	hi-lo-hi
	116578	D21262	Hs.75337	nucleolar phosphoprotein p130	hi-lo-hi
	130350	AA369601	Hs.239138	pre-B-cell colony-enhancing factor	hi-lo-hi
	101045	J05614		gb:Human proliferating cell nuclear anti	hi-lo-hi
	101544	M31169		gb:Human propionyl-CoA carboxylase beta-	hi-lo-hi
35	113674	NM_014214	Hs.5753	inositol(myo)-1(or 4)-monophosphatase 2	hi-lo-hi
55	102260	AL039104	Hs.159557	karyopherin alpha 2 (RAG cohort 1, impor	hi-ło-hi
	100154	H60720	Hs.81892	KIAA0101 gene product	hi-lo-hi
					hi-lo-hi
	100199	BE562298	Hs.71827	KIAA0112 protein; homolog of yeast ribos	
40	100372	NM_014791	Hs.184339	KIAA0175 gene product	hì-lo-hi
40	100387	D83777	"Hs.75137	KIAA0193 gene product	hi-lo-hi
	131514	BE270734	"Hs.2795	lactate dehydrogenase A	hi-lo-hi
	102938	W27518	Hs.234489	lactate dehydrogenase B	hi-lo-hi
	105811	BE617695	Hs.286192	protein phosphatase 1, regulatory (inhib	hi-lo-hi
4.5	101013	BE300094	"Hs.227751	lectin, galactoside-binding, soluble, 1	hi-lo-hi
45	124148	BE300094	"Hs.227751	lectin, galactoside-binding, soluble, 1	hi-lo-hì
	102968	AU076611	Hs.154672	methylene tetrahydrofolate dehydrogenase	hi-lo-hi
	130149	AW067805	Hs.172665	methylenetetrahydrofolate dehydrogenase	hi-lo-hi
	114767	AI859865	Hs.154443	minichromosome maintenance deficient (S.	hi-lo-hì
	129168	Al132988	Hs.109052	chromosome 14 open reading frame 2	hi-lo-hi
50	105011	BE091926	Hs.16244	mitotic spindle coiled-coil related prot	hi-lo-hi
	103023	AW500470	Hs.117950	multifunctional polypeptide similar to S	hi-lo-hì
	102808	BE242818	"Hs.179606	nuclear RNA helicase, DECD variant of DE	hi-lo-hi
	318617	AW247252	Hs.75514	nucleoside phosphorylase	hi-lo-hi
	101568	M81740	Hs.75212	ornithine decarboxylase 1	hi-lo-hi
55	102076	BE299197	Hs.179665	cyclin-dependent kinase inhibitor 1A (p2	hi-lo-hi
	100202	BE294407	"Hs.99910	phosphofructokinase, platelet	hi-lo-hi
	101032	BE206854	Hs.46039	phosphoglycerate mutase 2 (muscle)	hi-lo-hi
	130553	AF062649	*Hs.252587	pituitary tumor-transforming 1	hi-lo-hi
	101626	M57399	Hs.44	pleiotrophin (heparin binding growth fac	hi-lo-hi
60	101992	X90725	Hs.77597	polo (Drosophia)-like kinase	hì-lo-hi
50				procollagen-lysine, 2-oxoglutarate 5-dio	hi-lo-hi
	132164	AI752235	Hs.41270		
	101396	BE267931	"Hs.78996	proliferating cell nuclear antigen	hi-lo-hi
	119018	AA631143	Hs.179809	ESTs	hi-lo-hi
<b>C E</b>	101840	AA236291	Hs.183583	serine (or cysteine) proteinase inhibito	hi-lo-hi
65	332640	BE568452	Hs.5101	protein regulator of cytokinesis 1	hi-lo-hi
	132543	BE568452	Hs.5101	protein regulator of cytokinesis 1	hi-to-hi
	101118	AA371931	"Hs.77422	proteolipid protein 2 (colonic epitheliu	hi-to-hi
	109166	AA219691	Hs.73625	RAB6 interacting, kinesin-like (rabkines	hi-lo-hi
~~	100830	AC004770	"Hs.4756	flap structure-specific endonuclease 1	hi-lo-hi
70	107059	BE614410	Hs.23044	RAD51 (S. cerevisiae) homolog (E coli Re	hi-lo-hi
	321693	AA227069	Hs.173737	ras-related C3 botulinum toxin substrate	hi-lo-hi
	101148	NM_002923	Hs.78944	regulator of G-protein signalling 2, 24k	hi-lo-hi
	130567	AA383092	Hs.1608	replication protein A3 (14kD)	hì-lo-hi
	103076	NM_001034	Hs.75319	ribonucleotide reductase M2 polypeptide	hi-lo-hi
75	103131	BE536069	Hs.2962	S100 calcium-binding protein P	hi-lo-hi
- <del>-</del>	102212	AW411491	Hs.75069	serine hydroxymethyltransferase 2 (miloc	hi-lo-hi
	104254	AW411425	Hs.180655	serine/threonine kinase 12	hi-lo-hi
	102748	BE018138	Hs.24447	sigma receptor (SR31747 binding protein	hi-lo-hi
	102012	BE259035	Hs.118400	singed (Drosophila)-like (sea urchin fas	hi-lo-hi
80	102522	BE250944	Hs.183556	solute carrier family 1 (neutral amino a	hi-lo-hi
00	132994	AA112748	Hs.279905	clone HQ0310 PRO0310p1	hi-lo-hi
	101971	Z49105	"Hs.289105	synovial sarcoma, X breakpoint 2	hi-lo-hi
	1013/1	6-TJ 10J	110.403103	Sylvator scroomer v meanbount 5	10-0-13
				100	

	126645	AA316181	Hs.61635	six transmembrane epithelial antigen of	hi-lo-hi
	103058	X57348	Hs.184510	stratifin	hi-lo-hi
	102632	U66618	Hs.250581	SWI/SNF related, matrix associated, acti	hi-lo-hi
	103269	AF230662	"Hs.289105	synovial sarcoma, X breakpoint 2	hi-lo-hi
5	128920	AA622037	Hs.166468	programmed cell death 5	hi-lo-hi
,	100114	X02308	Hs.82962	thymidylate synthelase	hi-lo-hi
					hi-lo-hi
	102846	BE264974	Hs.6566	thyroid hormone receptor interactor 13	
	131877	J04088	"Hs.156346	topoisomerase (DNA) II alpha (170kD)	hi-lo-hi
10	100866	U14134	Hs.75113	general transcription factor IIIA	hi-lo-hi
10	133893	A1434699	Hs.77356	transferrin receptor (p90, CD71)	hi-lo-hi
	130135	AA311426	"Hs.21635	tubulin, gamma 1	hi-lo-hi
	130287	AA479005	Hs.154036	turnor suppressing subtransferable candid	hi-lo-hi
	126180	L32977	Hs.3712	ubiquinol-cytochrome c reductase, Rieske	hi-lo-hi
	101536	NM_006002	Hs.77917	ubiquitin carboxyl-terminal esterase L3	hi-lo-hi
15	102687	NM_007019	"Hs.93002	ubiquitin carrier protein E2-C	hì-lo-hi
13					
	103556	Z19002	Hs.37096	zinc finger protein 145 (Kruppel-like, e	hi-lo-hi
	300022				hi-lo-hi-lo
	133015	AJ002744	Hs.246315	UDP-N-acetyl-alpha-D-galactosamine:potyp	hi-lo-hi-lo
••	129642	NM_001360	Hs.11806	7-dehydrocholesterol reductase	hi-lo-lo
20	134369	AF207664	Hs.8230	a disintegrin-like and metalloprotease (	hi-lo-lo
	300023			, ,	hi-lo-lo
	125183	AV660804	Hs.301417	AHNAK nucleoprotein (desmoyokin)	hi-lo-lo
	101766	M80899	"Hs.301417	AHNAK nucleoprotein (desmoyokin)	hi-lo-lo
				annexin A2	hi-lo-lo
25	133516	BE265133	"Hs.217493		
23	102146	AW162057	Hs.78629	ATPase, Na+/K+ transporting, beta 1 poly	hi-lo-lo
	318538	A1750979	Hs.74034	Homo sapiens clone 24651 mRNA sequence	hi-lo-lo
	103554	A1878826	Hs.323469	caveolin 1, caveolae protein, 22kD	hi-lo-lo
	329365			CH.X_hs gi[5868838	hi-lo-lo
	334282			CH22_FGENES.369_12	hi-lo-lo
30	334891			CH22_FGENES.452_5	hi-lo-lo
50				CH22_FGENES.499_5	hi-lo-lo
	335149				
	335682			CH22_FGENES.595_2	hi-lo-lo
	335756			CH22_FGENES.604_5	hi-lo-lo
0.5	303951	AW475081	Hs.172928	collagen, type I, alpha 1	hi-lo-lo
35	134421	AU077196	Hs.82985	collagen, type V, alpha 2	hi-lo-lo
	131101	BE387561	Hs.22981	DKFZP586M1523 protein	hi-lo-lo
	124153	AU077333	*Hs.160483	erythrocyte membrane protein band 7.2 (s	hi-lo-lo
	103328	AU077333	"Hs.160483	erythrocyte membrane protein band 7.2 (s	hi-lo-lo
			*Hs.306201	hypothetical protein DKFZp564O1278	hi-lo-lo
40	322035	AL137517			
40	301872	H84730	Hs.326391	ESTs, Highly similar to KIAA1437 protein	hi-lo-lo
	303820	AB037858	Hs.173484	hypothetical protein FLJ10337	hi-lo-lo
	304049	T58155		gb:yb98h03.s1 Stratagene lung (937210) H	hl-lo-lo
	304735	AA576453		gb:nm75h11.s1 NCI_CGAP_Co9 Homo sapiens	hi-lo-lo
	306999	AJ138628	Hs.308058	EST, Weakly similar to zinc finger prot	hi-lo-lo
45	128789	AW368576	Hs.139851	caveolin 2	hi-lo-lo
	132057	AB037858	Hs.173484	hypothelical protein FLJ10337	hi-lo-lo
			Hs.173484	hypothetical protein FLJ10337	hi-lo-lo
	114795	AB037858			
	104204	AK001691	Hs.57655	hypothetical protein FLJ10829	hi-lo-lo
50	105200	AA328102	Hs.24641	cytoskeleton associated protein 2	hi-lo-lo
50	105493	AL047586	Hs.10283	RNA binding motif protein 88	hi-lo-lo
	107977	AI188161	Hs.144627	ESTs	hi-ło-ło
	108880	AA766605	"Hs.47099	hypothetical protein FLJ21212	hi-lo-lo
	111157	AL109729	Hs.18948	ESTs, Highly similar to A31026 probable	hì-lo-lo
	116202	BE159395	Hs.87089	ESTs	hi-lo-lo
55	120689	AW134519	Hs.96125	ESTs	hi-lo-lo
55		AA446628	Hs.2799	cartilage linking protein 1	hi-lo-lo
	121847				
	124182	. Al637471	Hs.107801	ESTs :	hi-lo-lo
	128515	BE395085	Hs.10086	type I transmembrane protein Fn14	hi-lo-lo
<b>CO</b>	130466	W19744	Hs.180059	Homo sapiens cDNA FLJ20653 fis, clone KA	hi-lo-lo
60	131076	AA749230	Hs.22666	ESTs	hi-lo-lo
	131084	NM_017413	Hs.303084	apelin; peptide ligand for APJ receptor	hi-lo-lo
	134109	AA348031	Hs.7913	ESTs	hì-lo-lo
	300258	Al478933	Hs.188260	ESTs	hi-lo-lo
•	302767	H94900	Hs.17882	ESTs	hi-lo-lo
65			Hs.133159	ESTs, Weakly similar to PIHUSD salivary	
0.5	312391	R43707			hi-lo-lo
	312689	AW450461	Hs.203965	ESTs	hi-lo-lo
	315715	AJ284219	Hs.130749	ESTs ·	hi-lo-lo
	315843	AA679430	Hs.191897	ESTs	hi-lo-lo
	322447	A1735759	Hs.52620	integrin, beta 8	hì-lo-lo
70	322826	AI807883	Hs.201771	ESTs	hi-lo-lo
-	324867	A1624707	"Hs.5921	Homo sapiens cDNA: FLJ21592 fis, clone C	hi-lo-lo
	331336	AA287450	Hs.93842	Homo sapiens cDNA: FLJ22554 fis, done	hi-to-to
				ESTs	hi-lo-lo
	331353	AA953006	Hs.88143		
75	133063	A1654133	Hs.30212	thyroid receptor interacting protein 15	hi-lo-lo
75	311034	BE567130	Hs.311389	ESTs, Moderately similar to PT0375 natur	hi-lo-lo
	108647	BE546947	Hs.44276	homeo box C10	hi-lo-lo
	124955	AA376768	°Hs.324841	hypothetical protein FLJ22622	hi-lo-lo
	113923	AW953484	Hs.3849	hypothetical protein FLJ22041 similar to	hi-lo-lo
	310557	Al431798	Hs.164192	ESTs, Weakly similar to Y161_HUMAN HYPOT	hi-lo-lo
80	302943	AI581344	Hs.127812	ESTs, Weakly similar to T17330 hypotheti	hi-lo-lo
30					
	128453	X02761	"Hs.287820	fibronectin 1	hi-ko-lo
	305232	AA670052	Hs.169476	glyceraldehyde-3-phosphate dehydrogenase	hi-lo-lo
				101	

	447040		D1 45445		
	117642	U55184	"Hs.154145	hypothetical protein FLJ11585	hi-lo-lo
	115881 133666	NM_005756 U56725	Hs.184942 Hs.75452	G protein-coupled receptor 64 heat shock 70kD protein 2	hi-lo-lo hi-lo-lo
	103262	X78565	Hs.289114	hexabrachion (tenascin C, cytotactin)	hi-lo-lo
5	100793	S69027	113.203114	gb:HOX C6=class I homeodomain (fragment	hi-lo-lo
•	102289	U32114		gon to to the transfer and the great	hi-lo-lo
	319109	Z45662	Hs.90797	Homo sapiens clone 23620 mRNA sequence	hi-lo-lo
	116357	AF052107	Hs.90797	Homo sapiens clone 23620 mRNA sequence	hi-lo-lo
10	101497	W05150	"Hs.37034	homeo box A5	hi-lo-lo
10	105508	AA173942	Hs.326416	Homo sapiens mRNA; cDNA DKFZp564H1916 (f	hi-lo-lo
	302290	AA179949	Hs.175563	Homo sapiens mRNA; cDNA DKFZp564N0763 (f	hi-lo-lo
	102838	R34657	Hs.80658	uncoupling protein 2 (mitochondrial, pro	hi-lo-lo
	100235	D29954 NM 002206	Hs.13421 Hs.74369	KIAA0056 protein	hi-lo-lo hi-lo-lo
15	133507 125573	Al351642	Hs.182241	integrin, alpha 7 interferon induced transmembrane protein	hi-lo-lo
13	103059	X57351	Hs.174195	interferon induced transmembrane protein	hi-lo-lo
	330415	D83777	*Hs.75137	KIAA0193 gene product	hi-lo-lo
	303054	BE265848	Hs.289080	colon cancer-associated protein Mic1	hi-lo-lo
	133579	X75346	Hs.75074	mitogen-activated protein kinase-activat	hi-lo-lo
20	100528	BE386801	Hs.21858	trinucleotide repeat containing 3	hi-lo-lo
	107480	AF001691	Hs.74304	periplakin	hi-lo-lo
	133050	X73424	Hs.63788	propionyl Coenzyme A carboxylase, beta p	hi-lo-lo
	133061	AI186431	Hs.296638	prostate differentiation factor	hi-lo-lo
25	106390	AJ297436	Hs.20166	prostate stem cell antigen	hi-lo-lo
25	302124	AA676403	Hs.145078	regulator of differentiation (in S. pomb	hi-lo-lo
	129823	X00949	"Hs.105314	relaxin 1 (H1)	hi-lo-lo
	134444 103240	BE184455 U81961	"Hs.251754 Hs.2794	secretory leukocyte protease inhibitor ( sodium channel, nonvoltage-gated 1 alpha	hi-lo-lo hi-lo-lo
	115761	AA366037	Hs.90911	solute carrier family 16 (monocarboxylic	hi-lo-lo
30	321412	AI674383	Hs.22891	solute carrier family 7 (cationic amino	hi-lo-lo
50	126487	AA283809	Hs.184601	solute carrier family 7 (cationic amino	hi-lo-lo
	101759	M80244	Hs.184601	solute carrier family 7 (cationic amino	hi-lo-lo
	112941	AW163034	Hs.6467	synaptogyrin 3	hi-lo-lo
	134351	BE272506	"Hs.82109	syndecan 1	hi-lo-lo
35	125924	BE272506	"Hs.82109	syndecan 1	hì-lo-lo
	130982	AA033627	Hs.21858	trinucleotide repeat containing 3	hi-lo-lo
	133473	AW301993	Hs.73980	troponin T1, skeletal, slow	hi-lo-lo
	101042	T46839	"Hs.10319	UDP glycosyltransferase 2 family, polype	hi-lo-lo
40	129565	X77777	Hs.198726	vasoactive intestinal peptide receptor 1	hi-lo-lo
40	102992 106868	M85430 BE185536	"Hs.155191 Hs.300816	villin 2 (ezrin) Homo sapiens mRNA; cDNA DKFZp564I172 (fr	hi-lo-lo lo-hi lo
	132618	AL050025	"Hs.279916	hypothetical protein FLJ20151	lo-hi-hi
	100187	D17793	*Hs.78183	aldo-keto reductase family 1, member C3	lo-hi-hi
	116334	AL038450	Hs.48948	ATP2C1 calcium transport ATPase, same as	lo-hi-hi
45	134454	NM_013230	Hs.286124	CD24 antigen (small cell lung carcinoma	lo-hì-hi
	302067	BE542706	Hs.222399	CEGP1 protein	lo-hi-hi
	105500	AW602166	Hs.222399	CEGP1 protein	lo-hi-hi
	100732	AA557660	°Hs.76152	decarin	lo-hi-hi
50	129265	AA530892	Hs.171695	dual specificity phosphatase 1	lo-hi-hi
50	117789	N48294	Hs.46850	EST	lo-hi-hi
	330786	BE379594	"Hs.49136	ESTs, Moderately similar to ALU7_HUMAN A hypothetical protein FLJ10890	lo-hi-hi lo-hi-hi
	319808 303502	T58960 BE174240	Hs.17283	gb:QV1-HT0573-290200-092-f06 HT0573 Homo	lo-hi-hi
	116780	H22566	"Hs.30098	ESTs	lo-hi-hi
55	104189	AB040927	Hs.301804	KIAA1494 protein	lo-hi-hi
-	105588	L43821	Hs.80261	enhancer of filamentation 1 (cas-like do	lo-hi-hi
	105731	AA834664	Hs.29131	nuclear receptor coactivator 2	lo-hi-hi
	105772	H57111	Hs.221132	ESTs	lo-hì-hi
<b>60</b>	105794	H24530	Hs.273294	hypothetical protein FLJ20069	lo-hi-hi
60	113098	N77737	Hs.8349	Apobec-1 complementation factor; APOBEC-	lo-hi-hi
	113803	AW880709	*Hs.283683	chromosome 8 open reading frame 4	lo-hi-hi
	114530	AA601038	Hs.191797	ESTs	lo-hi-hi
	116188	AA468183	Hs.184598	Homo sapiens cDNA: FLJ23241 fis, clone C	lo-hi-hi lo-hi-hi
65	117330 117701	AI904095 BE063921	Hs.43423 Hs.295971	ESTs ESTs	lo-hi-hi
05	120911	Al189754	Hs.144330	ESTs	lo-hi-hi
	124083	AW195237	Hs.7734	hypothetical protein FLJ22174	lo-hì-hi
	124690	AW883529	Hs.173830	ESTs	lo-hi-hi
	130796	AA088809	Hs. 19525	hypothetical protein FLJ22794	lo-hi-hi
70	131524	AB040927	Hs.301804	KIAA1494 protein	lo-hi-hì
	132116	AW960474	Hs.40289	ESTs	lo-hi-hi
	132442	AW970859	Hs.313503	ESTs	lo-hi-hi
	310219	AI221087	Hs.147761	ESTs	lo-hi-hi
75	310598	Al439136	Hs.140546	ESTs	lo-hi-hi
75	310884	AW014684	Hs.232189	ESTS ESTS Markly similar to SMM1 HILMAN SURVI	lo-hi-hi lo bi bi
	311587	A1828254	Hs.271019	ESTs, Weakly similar to SMN1_HUMAN SURVI Homo sapiens cDNA FLJ12028 fis, done HE	lo-hi-hi lo-hi-hi
	312240 312803	R36475 AA677934	Hs.24321 Hs.117864	ESTs	lo-hi-hi
	312803 314219	AA677934 AA262331	Hs.48376	Homo sapiens clone HB-2 mRNA sequence	lo-hi-hi
80	315052	AA876910	Hs.134427	ESTs	lo-hi-hi
	331919	AA446869	Hs.119316	ESTs	lo-hi-hi
	133240	AK001489	Hs.242894	ADP-ribosylation factor-like 1	lo-hi-hi
				100	

	134006	Z45957	Hs.7837	G-protein-coupled receptor induced prote	lo-hi-hi
	124847	W07701	"Hs.304177	Homo sapiens clone FLB8503 PRO2286 mRNA	lo-hi-hi
	129087	Al348027	Hs.108557	Homo sapiens clone PP1057 unknown mRNA	lo-hi-hi
	131762	AA744902	"Hs.107767	hypothetical protein PRO1489	
5					lo-hi-hi
,	129000	AA744902	"Hs.107767	hypothetical protein PRO1489	lo-hi-hi
	105713	Al122843	"Hs.184319	ESTs, Weakly similar to KIAA1006 protein	lo-hi-hi
	118475	N66845		gb:za46c11.s1 Soares fetal liver spleen	lo-hi-hi
	118381	N64513	Hs.48994	ESTs, Weakly similar to AF151800 1 CGI-4	lo-hi-hi
	105057	AA134233		gb:zo20f10.s1 Stratagene colon (937204)	lo-hi-hi
10	131507	Al826268	Hs.27769	ESTs, Wealty similar to MCAT_HUMAN MITOC	lo-hi-hì
	124970	BE272862	Hs.106534	hypothetical protein FLJ22625	
	130094	NM_001471	*Hs.167017		lo-hi-hi
	302357			gamma-aminobutyric acid (GABA) B recepto	lo-hi-hi
		X03178	Hs.198246	group-specific component (vitamin D bind	lo-hi-hi
1.5	113231	AA278583	Hs.180737	Homo sapiens clone 23664 and 23905 mRNA	lo-hi-hi
15	111923	BE383234	Hs.25925	Homo sapiens clone 23860 mRNA sequence	lo-hi-hi
	128530	Al932995	Hs.183475	Homo sapiens clone 25061 mRNA sequence	lo-hi-hi
	128987	A1339046	Hs.107637	hypothetical protein FLJ12806	lo-hi-hi
	315368	AB037745	Hs.104696	KIAA1324 protein	lo-hi-hi
	133944	AW068579	Hs.7780	Homo sapiens mRNA; cDNA DKFZp564A072 (fr	
20					lo-hi-hi
20	115084	BE383668	"Hs.42484	hypothetical protein FLJ10618	lo-hi-hi
	132883	AA373314	Hs.5897	Homo sapiens mRNA; cDNA DKFZp586P1622 (f	lo-hi-hi
	109623	AW207385	Hs.295901	KIAA0493 protein	lo-hi-hi
	130577	M69241	"Hs.162	insulin-like growth factor binding prote	lo-hi-hi
	101889	AF188747	"Hs.181350	kallikrein 2, prostatic	lo-hi-hi
25	130336	AA535210	"Hs.171995	kallikrein 3, (prostate specific antigen	lo-hi-hi
	128180	AW949068	Hs.171995	kallikrein 3, (prostate specific antigen	
	134921				lo-hi-hi
		AL137491	Hs.125511	Homo sapiens mRNA; cDNA DKFZp434P1530 (f	lo-hi-hi
	302385	AJ224172	Hs.204096	lipophilin B (uteroglobin family member)	lo-hi-hi
20	117921	AA021459	Hs.306480	Homo sapiens mRNA; cDNA DKFZp761E2112 (f	lo-hi-hi
30	101701	NM_002436	Hs.1861	membrane protein, palmitoylated 1 (55kD)	lo-hi-hì
	130356	AF127577	Hs.155017	nuclear receptor interacting protein 1	lo-hi-hì
	101763	AB001914	Hs.170414	paired basic amino acid cleaving system	lo-hì-hi
	130342	U81802	Hs.154846	phosphatidylinositol 4-kinase, catalytic	
	130760	AW379130			lo-hi-hi
35			Hs.18953	phosphodiesterase 9A	lo-hi-hi
33	101461	N98569	Hs.76422	phospholipase A2, group IIA (platelets,	lo-hi-hi
	134032	NM_005025	Hs.78589	serine (or cysteine) proteinase inhibito	lo-hi-hi
	303762	AF034799	Hs.30881	protein tyrosine phosphatase, receptor t	lo-hi-hi
	110932	AA021459	Hs.306480	Homo sapiens mRNA; cDNA DKFZp761E2112 (f	lo-hi-hi
	135192	U83993	Hs.321709	purinergic receptor P2X, ligand-gated to	lo-hi-hi
40	133886	U97276	Hs.77266	quiescin Q6	lo-hi-hi
. •	134142	BE244053	Hs.79362	retinoblastoma-like 2 (p130)	
					lo-hi-hi
	100877	X80821	Hs.302177	H.sapiens mRNA for ribosomal protein L18	lo-hi-hi
	133534	AU077115	Hs.201675	RNA binding motif protein 5	lo-hi-hi
10	133011	NM_006379	Hs.171921	sema domain, immunoglobulin domain (lg),	lo-hi-hi
45	132160	W26406	Hs.295923	seven in absentia (Drosophila) homolog 1	lo-hi-hi
	103110	X62822	Hs.2554	sialyltransferase 1 (beta-galactoside al	lo-hi-hi
	130173	U38847	Hs.151518	TAR (HIV) RNA-binding protein 1	lo-hi-hi
	127435	X69086	"Hs.286161	Homo sapiens cDNA FLJ13613 fis, clone PL	lo-hi-hi
	110520	N54069	Hs.4082	lectin, galactoside-binding, soluble, 8	
50	114660	AA071383	143.4002		lo-hi-hi
50			11-000007	gb:zm61d05.r1 Stratagene fibroblast (937	lo-hi-hi
	330541	NM_002038	Hs.265827	interferon, alpha-inducible protein (clo	lo-hi-lo
	101486	AA506324	Hs.1852	acid phosphalase, prostate	lo-hi-lo
	332386	NM_000481	Hs.102	aminomethyltransferase (glycine cleavage	lo-hi-lo
	100569	AA535210	"Hs.171995	kallikrein 3, (prostate specific antigen	lo-hi-lo
55	134738	AU076801	Hs.89436	cadherin 17, Ll cadherin (liver-intestin	lo-hi-lo
	103119	X63629	Hs.2877	cadherin 3, type 1, P-cadherin (placenta	
	302892	AW176909	Hs.42346		lo-hi-lo
				calcineurin-binding protein calsarcin-1	lo-hi-lo
	105402	AB014680	Hs.8786	carbohydrate (chondroitin 6/keratan) sul	lo-hi-lo
60	102976	AU077174	"Hs.288181	cathepsin H	to-hi-to
60	101793	W01076	"Hs.119663	CD59 antigen p18-20 (antigen identified	lo-hi-lo
	129890	Al868872	"Hs.282804	Homo sapiens cDNA: FLJ22704 fis, clone H	lo-hi-to
	328164			CH.06_hs gij5868068	lo-hi-lo
	328648			CH.07_hs gi 6004473	lo-hi-la
	330032			CH.16_p2 glj6682596	lo-hi-lo
65	330033			CH.16_p2 gij6682596	
05	326816				to-hi-lo
				CH.20_hs gi]6552458	lo-hi-lo
	337603			CH22_C20H12.GENSCAN.16-2	lo-hi-to
	338561			CH22_EM:AC005500.GENSCAN.421-5	lo-hi-lo
	338562			CH22_EM:AC005500.GENSCAN,421-6	lo-hi-lo
70	333743			CH22_FGENES.264_1	lo-hi-lo
	333845			CH22_FGENES.290_3	lo-hi-lo
	333849			CH22_FGENES.290_8	
					lo-hi-lo
	334221			CH22_FGENES.360_1	lo-hi-lo
75	334222			CH22_FGENES.360_3	lo-hi-lo
75	334578			CH22_FGENES.406_1	lo-hi-lo
	336662			CH22_FGENES.41-1	lo-hi-lo
	336684			CH22_FGENES.46-1	lo-hi-lo
	335289			CH22_FGENES.527_2	lo-hi-lo
	335290				
80				CH22_FGENES.527_3	lo-hi-lo
OU.	335293			CH22_FGENES.527_6	lo-hi-lo
	337182			CH22_FGENES.570-2	lo-hi-lo
	335809			CH22_FGENES.617_6 (same as BFH4)	lo-hi-lo
				102	

	335810			CH22_FGENES.617_7	lo-hi-lo
	335824			CH22_FGENES.619_11 (same as BFH5)	lo-hi-lo
					lo-hi-lo
	336054			CH22_FGENES.683_3	lo-hi-lo
_	333124			CH22_FGENES.81_8	
5	332340	AP000692	Hs.129781	chromosome 21 open reading frame 5	lo-hi-lo
	130380	AI949359	Hs.143600	type II Golgi membrane protein	lo-hi-lo
	102962	R50032	Hs.159263	collagen, type VI, alpha 2	to-hi-lo
	331306	AF102546	Hs.63931	dachshund (Drosophila) homolog	to-hi-lo
	319408	AA448090	Hs.87359	ESTs, Highly similar to RB18 MOUSE RAS-R	lo-hi-lo
10	312197	T96203		gb:ye48b07.r1 Soares fetal liver spleen	lo-hi-lo
10	312405	AI523875		gb:tg97d04.x1 NCI_CGAP_CLL1 Homo sapiens	lo-hi-lo
	312939	AA495930	Hs.24444	Homo sapiens cDNA: FLJ22165 fis, clone H	lo-hi-lo
					lo-hi-lo
	313475	AA010200	Hs.175551	ESTs	
1.5	313624	AA525775	Hs.292523	ESTs	lo-hi-lo
15	316897	AA838114	Hs.221612	ESTs	lo-hi-lo
	317850	Al681545	Hs.152982	hypothetical protein FLJ13117	lo-hi-lo
	318541	T30290	Hs.107515	ESTs	ło-hi-lo
	321325	AB033100	Hs.300646	KIAA protein (similar to mouse paladin)	lo-hi-lo
	321696	AA628791	Hs.76228	amplified in osteosarcoma	lo-hi-lo
20	322189	H65014	10.10220	gb:yu66f10.r1 Weizmann Olfactory Epithel	lo-hi-lo
20			Hs.137306		lo-hi-lo
	322463	Al242754	HS. 13/300	ESTs	
	322540	R76593		gb:yi60c11.r1 Soares placenta Nb2HP Homo	lo-hi-lo
	323131	AK002088	Hs.270124	Homo sapiens cDNA FLJ11226 fis, clone PL	lo-hi-lo
0.5	323243	W47525	Hs.110771	Horno sapiens cDNA: FLJ21904 fis, clone H	lo-hi-lo
25	323591	AA301270		gb:EST14192 Testis tumor Homo sapiens cD	lo-hi-lo
	323753	AK002161	Hs.70266	yeast Sec31p homolog	lo-hi-lo
	323835	AL042005	Hs.1117	tripeptidyl peptidase II	lo-hi-lo
	323926	AA354572		gb:EST62857 Jurkat T-cells V Homo sapien	lo-hi-lo
	324047	AI433357	"Hs.271340	ESTs	to-hi-lo
30			H5.2/ 1340		lo-hi-lo
20	324330	AA884766		gb:am20a10.s1 Soares_NFL_T_GBC_S1 Homo s	
	324753	AA612626	Hs.144871	Homo sapiens cDNA FLJ13752 fis, clone PL	lo-hi-lo
	300702	AA075481	Hs.111334	ferritin, light polypeptide	lo-hi-lo
	301712	BE083080	Hs.274323	Homo sapiens, Similar to sialyttransfera	lo-hi-lo
	302380	AA325633	Hs.136102	KIAA0853 protein	lo-hi-lo
35	302970	W05608	Hs.312679	EST	lo-hi-lo
-	303187	AA115962	Hs.323423	ESTs, Moderately similar to B Chain B,	to-hi-lo
	303194	AA082000	113.020720	gb:zn26f07.r1 Stratagene neuroepithelium	lo-hi-lo
			Un 272572		lo-hi-lo
		· AA782347	Hs.272572	hemoglobin, alpha 2	
40	304263	AA062837		gb:zm05b11.s1 Stratagene corneal stroma	lo-hi-lo
40	304275	AA070605		gb:zm53h09.s1 Stratagene fibroblast (937	lo-hi-lo
	304309	AA112147		gb:zm64c06.s1 Stratagene fibroblast (937	lo-hi-lo
	305503	AA759177	Hs.298148	ESTs, Weakly similar to KIAA0565 protei	lo-hi-lo
	308615	AK000142	Hs.101774	hypothetical protein FLJ23045	lo-hi-lo
	309390	AW080585		gb:xc33f09.x1 NCI_CGAP_Co18 Homo sapiens	to-hi-lo
45	104667	Al239923	Hs.30098	ESTs	lo-hi-lo
	310014	D60745	Hs.25925	Homo sapiens clone 23860 mRNA sequence	lo-hi-lo
	318814	W07361	Hs.22545	Homo sapiens cDNA FLJ12935 fis, clone NT	lo-hi-lo
			Hs.47191	ESTs	lo-hi-lo
	321896	C04863			lo-hi-lo
50	331661	W52448	Hs.56147	ESTs	
50	332120	AA609684	Hs.112748	Homo sapiens cDNA: FLJ21543 fis, clone C	lo-hi-lo
	332256	AW975028	Hs.102754	ESTs	lo-hi-lo
	107252	D60745	Hs.25925	Homo sapiens clone 23860 mRNA sequence	lo-hi-lo
	112068	A1264847	Hs.22545	Homo sapiens cDNA FLJ12935 fis, clone NT	lo-hi-lo
	117929	N51075	Hs.47191	ESTs	lo-hi-to
55	119637	W52448	Hs.56147	ESTs	lo-hi-lo
	123712	AA609684	Hs.112748	Homo sapiens cDNA: FLJ21543 fis, clone C	lo-hi-lo
	124560	AW975028	Hs.102754	ESTs	lo-hi-io
	105039	AA907305	Hs.36475	ESTS	to-hi-lo
60	105271	AA807881	Hs.25329	ESTs	lo-hi-lo
60	106689	AW296584	Hs.293782	ESTs	lo-hi-lo
	106849	AL137281	Hs.17110	Homo sapiens mRNA; cDNA DKFZp434C2016 (f	lo-hi-lo
	107071	AW385224	Hs.35198	ectonucleotide pyrophosphatase/phosphodi	lo-hi-lo
	108218	W57550	Hs.301526	hypothetical protein FLJ13181	lo-hi-lo
	110930	BE242691	Hs.14947	ESTs, Weakly similar to ALU1_HUMAN ALU S	lo-hi-lo
65	112098	R44714	Hs.106795	Homo sapiens cDNA FLJ13136 fis, clone NT	lo-hi-lo
UJ					lo-hi-lo
	112170	BE246743	Hs.288529	hypothetical protein FLJ22635	lo-hi-lo
	112902	AL035633	"Hs.129190	Human DNA sequence from clone RP5-1046G1	
	114877	AW024162	Hs.205125	ESTs	lo-hi-lo
70	116312	BE379794	Hs.65403	hypothetical protein	lo-hi-to
70	116739	H01463	Hs.93534	ESTs	lo-hi-lo
	119267	AA064970	Hs.118145	ESTs	lo-hi-lo
	120570	AA280679	Hs.271445	ESTs, Weakly similar to ALU1_HUMAN ALU	lo-hi-lo
	121176	AL121523	Hs.97774	ESTs	lo-hi-lo
	123360	AA532718	Hs.178604	ESTs	lo-hi-lo
75					lo-hi-lo
75	123974	NM_015678	Hs.3821	neurobeachin	
	124777	R41933		gb:yg04f09.s1 Soares infant brain 1NIB H	lo-hi-lo
	128046	AA873285	•	gb:oh68h05.s1 NCI_CGAP_Kid5 Homo sapiens	lo-hi-lo
	128666	AA808466	Hs.103395	hypothetical protein FLJ14146	lo-hi-lo
	130639	AI557212	"Hs.17132	ESTs	lo-hi-lo
80	130693	R68537	Hs.17962	ESTs	lo-hi-lo
00	131756	AA443966	Hs.31595	ESTs	lo-hi-lo
		AA503020	Hs.36563	hypothetical protein FLJ22418	lo-hi-lo
	131985	MANUSTER	113.00303	пурована риман с мести	טרוורט

	132932	AW118826	Hs.6093	Homo sapiens cDNA: FLJ22783 fis, clone K	lo-hi-lo
	134696	BE326276	°Hs.8861	ESTs	lo-hi-lo
	300967	AA565209	Hs.269439	ESTs	lo-hi-lo
	301182	AW291411	Hs.192531	ESTs, Weakly similar to S00754 zinc fing	lo-hi-lo
5	302595	Al699372	Hs.193247	Homo sapiens mRNA; cDNA DKFZp434A171 (fr	lo-hi-lo
-	303132	Al929819	Hs.4055	chromosome 21 open reading frame 50	to-hi-lo
	303506	AA340605	Hs.105887	ESTs, Weakly similar to Homolog of rat Z	lo-tri-lo
	303654	BE246743	Hs.288529	hypothetical protein FLJ22635	lo-hi-lo
	310026	AA278233	Hs.100691	ESTs	lo-hi-lo
10	310056	AI253072	Hs.145383	ESTs	io-hi-lo
10	310353	AI261700	Hs.145544	ESTs	io-hi-lo
	310371	Al262584	Hs.145575	ESTs	lo-hi-lo
	310430	Al670843	Hs.200257	ESTs	lo-hi-lo
	310438	AW022192	Hs.200197	ESTs	lo-hi-lo
15	310455	AI277603	Hs.145990	ESTs	lo-hi-lo
10	310787	AW262580	Hs.147674	KIAA1621 protein	lo-hi-lo
	311067	AI587332	Hs.209115	ESTs	lo-hi-lo
	311422	F00677	Hs.101316	ESTs	lo-hi-lo
	311465	Al758660	Hs.206132	ESTs	lo-hi-lo
20	312073	AA682393	Hs.119237	ESTs	lo-hi-lo
	312105	T81819	Hs.302251	ESTs	lo-tri-lo
	312108	T82331	*Hs.127453	ESTs	lo-hi-lo
	312292	AW450103	Hs.151124	ESTs	lo-hi-lo
	312313	AW293341	Hs.122505	ESTs, Weakly similar to 138022 hypotheti	lo-hi-lo
25	312600	AW970985	Hs.290853	ESTs	lo-hi-lo
	312800	Al248774	Hs.126707	hypothetical protein FLJ11457	lo-hi-lo
	312821	AA699325	Hs.269880	ESTs	lo-hi-lo
	313097	Al676164	Hs.204339	ESTs	lo-hi-lo
	313166	AI801098	Hs.151500	ESTs	lo-hi-lo
30	313179	AA927670	Hs.131704	ESTs	lo-hi-lo
- •	313280	AW960454	Hs.222830	ESTs	lo-hi-lo
	313689	Al608810	Hs.193288	ESTs	lo-hi-lo
	314146	AI827237	Hs.282884	ESTs	lo-hi-lo
	314305	Al280112	Hs.125232	Homo sapiens cDNA FLJ13266 fis, clone OV	lo-hi-lo
35	314456	Al867931	Hs.164595	ESTs	lo-hi-lo
	314465	AA602917	Hs.156974	ESTs	lo-hi-lo
	314881	A1095087	Hs.152299	ESTs, Moderately similar to ALU5_HUMAN A	lo-hi-to
	314916	AA548906	Hs.122244	ESTs	lo-hi-to
	315043	AA806538	Hs.130732	KIAA1575 protein	lo-hi-to
40	315074	AA828284	Hs.136729	Homo sapiens cDNA: FLJ21348 fis, clone C	lo-hi-lo
	315214	AI915927	Hs.34771	ESTs	lo-hi-lo
	315344	AW292176	Hs.245834	ESTs	lo-hi-lo
	315353	Al373949	Hs.279610	hypothetical protein FLJ10493	io-hi-lo
4.5	315439	T78413	Hs.293696	ESTs	io-hi-lo
45	315528	R37257	Hs.184780	ESTs	lo-hi-lo
	315720	AA292998	Hs.163900	ESTs	lo-hi-lo
	315772	AW515373	Hs.271249	Homo sapiens cDNA FLJ13580 fis, clone PL	lo-hi-lo
	315841	AW136397	Hs.247572	ESTs	lo-hi-lo
50	316042	AI469960	Hs.170698	ESTs	lo-hì-lo
50	316244	A1640761	Hs.224988	ESTs	lo-hi-lo
	316345	AW139408	Hs.152940	ESTs	lo-hi-lo
	316625	BE540090	Hs. 122156	ESTs	lo-hi-lo
	316738	AA889055	Hs.123468	ESTs	lo-tri-lo
55	316868	AI660898	Hs. 195602	ESTs	lo-hì-lo
55	316905	AW138241	Hs.210846	ESTs	lo-hì-lo
	317224	X73608	"Hs.93029	sparc/osteonectin, cwcv and kazal-like d	lo-hi-lo
	317275	AI809444	Hs.202108	ESTs	io-hi-lo
	317404	AI806867	Hs.126594	ESTs	lo-hi-lo
60	317488	AW071851	Hs.130628	ESTs	lo-hi-lo
UU	317916	AI565071	Hs.159983	ESTs	lo-hi-lo
	317939	Al986208	Hs.244760	ESTS	lo-hi-lo
	318486	T23514	11- 40000	gb:seq3329 1-NIB Homo sapiens cDNA clone	lo-hi-lo
	319897	N46574	Hs.43838	ESTs	lo-hi-lo
65	320654	AI160015	Hs.118112	ESTs	lo-hi-lo
05	320697	N62937	Hs.269109	ESTs	lo-hi-lo
	320787	AW088363	Hs.246240	ESTs	lo-hi-lo
	321023	AW294316	Hs.125608	ESTs	lo-hi-lo
	321899	AW972832	Hs.29468	ESTs	lo-hi-lo
70	322939 323045	AA101697 AA148950	Hs.211270	ESTs ESTs	lo-hi-lo
, 0	323045	AJ902456	Hs.188836	ESTs ESTs	lo-hi-lo
	323262	AL133990	Hs.210761	ESTs ECT:	lo-hi-lo
	323202 323410	AW118683	Hs.190642	ESTs ESTs	lo-hi-lo
	323645	AW445014	Hs.154150 He 197746	ESTs ESTs	lo-hi-lo
75	324598	AW972227	Hs.197746	Homo sapiens cDNA: FLJ22765 fis, clone K	lo-hi-lo
, 5	324666	T78413	Hs.163986 Hs.293696	ESTs	lo-hi-lo
	324674	AA541323	Hs.115831	ESTS	lo-hi-lo
	324713	A1093930	"Hs.313466	ESTS	lo-hi-lo
	324790	Al334367	Hs.159337	ESTs	lo-hi-lo lo-hi-lo
80	324804	AI692552	100.100001	gb:wd73f12.x1 NCI_CGAP_Lu24 Homo sapiens	lo-hi-lo
- •	330728	AI905520	Hs.29672	ESTs	lo-hi-lo
	330760	H04588	Hs.30469	ESTs	lo-hi-lo
				= :	11170

	330776	AW953805	Hs.21887	ESTs	lo-hi-lo
	330824	AB037732	Hs.61441	KIAA1311 protein	lo-hi-lo
	331028	A1539652	Hs.28338	KIAA1546 protein	lo-hi-lo
_	331046	N66563	Hs.191358	ESTs	lo-hi-lo
5	331050	BE007967	Hs.155795	ESTs	lo-hi-lo
	331053	AI949841	Hs.183146	ESTs, Moderately similar to ALU1_HUMAN A	lo-hi-lo
	331180	R44692	Hs.6640	Human DNA sequence from PAC 75N13 on chr	lo-hi-lo
	331313	AA761094	"Hs.80618	hypothetical protein	lo-hi-lo
10	331337	N74392	Hs.50495	ESTs	lo-hi-lo
10	331393	AW976438	°Hs.17428	RBP1-like protein	lo-hi-lo
	331432	AA262451	Hs.38485	ESTs	lo-hi-lo
	331517	AA765603	Hs.180877	H3 histone, family 3B (H3.3B)	lo-hi-lo
	331686	AW474960	Hs.182258	ESTs	lo-hi-lo
16	332002	A1579909	Hs.105104	ESTs	lo-hi-lo
15	332043	AA371307	Hs.125056	ESTs	lo-hi-lo
	332265	AW770320	Hs.222413	ESTs	to-hi-lo
	332314	R41396	Hs.101774	hypothetical protein FLJ23045	to-hi-lo
	131517	AB037789	Hs.263395	sema domain, transmembrane domain (TM),	lo-hi-lo
00	315352	AA604799	Hs.136528	ESTs, Moderately similar to ALU1_HUMAN A	lo-hi-la
20	315498	AA628539	Hs.116252	ESTs, Moderately similar to ALU1_HUMAN A	lo-hi-lo
	321489	Al459177	Hs.172759	ESTs, Moderately similar to ALU7_HUMAN A	lo-hi-lo
	106099	NM_012068	Hs.9754	activating transcription factor 5	lo-hi-lo
	105726	NM_012068	Hs.9754	activating transcription factor 5	lo-hi-lo
25	319926	Al820719	Hs.154662	DnaJ (Hsp40) homolog, subfamily A, membe	lo-hi-lo
25	314915	AI673735	Hs.187748	ESTs, Weakly similar to ALU1_HUMAN ALU S	lo-hi-lo
	315198	Al741506	Hs.186753	ESTs, Weakly similar to ALU1_HUMAN ALU S	lo-hi-lo
	324302	AW972771	Hs.292471	ESTs, Weakly similar to ALU1_HUMAN ALU S	lo-hi-lo
	331341	BE541042	"Hs.23240	Homo sapiens cDNA FLJ13496 fis, clone PL	lo-hi-lo
-	113783	AL359588	Hs.7041	hypothetical protein DKFZp762B226	lo-hi-lo
30	313552	AI889208	Hs.17283	hypothetical protein FLJ10890	lo-hi-lo
	103989	AA315993	Hs.105484	Homo saplens regenerating gene type IV m	lo-hi-lo
	331492	AK001114	Hs.53913	hypothetical protein FLJ10252	lo-hi-lo
	110837	H03109	Hs.108920	HTO18 protein	lo-hi-lo
25	330814	A1955040	Hs.265398	ESTs, Weakly similar to transformation-r	lo-hi-lo
35	312226	AA315703	Hs.199993	ESTs	lo-hi-lo
	102034	Al903474	Hs.230	fibromodulin	lo-hi-lo
	134671	BE263255	Hs.302749	FK506-binding protein 9 (63 kD)	lo-hi-lo
	131083	Y09763	Hs.22785	gamma-aminobutyric acid (GABA) A recepto	io-hi-lo
40	309575	AW168096	Hs.169476	glyceraldehyde-3-phosphate dehydrogenase	lo-hi-lo
40	134332	D86962	Hs.81875	growth factor receptor-bound protein 10	lo-hi-lo
	132904	NM_005518	Hs.59889	3-hydroxy-3-methylglutaryl-Coenzyme A sy	lo-hi-lo
	302910	N77976	Hs.251577	hemoglobin, alpha 1	lo-hi-lo
	133731	N71725	"Hs.272572	hemoglobin, alpha 2	lo-hi-lo
4.5	303297	AF070623	Hs.13423	Homo sapiens clone 24468 mRNA sequence	lo-hi-lo
45	108732	AA258888	Hs.107476	ATP synthase, H+ transporting, mitochond	lo-hi-lo
	108731	AA258888	Hs.107476	ATP synthase, H+ transporting, mitochond	lo-hi-lo
	302123	AB013452	Hs.144931	ATPase, aminophospholipid transporter (A	lo-hi-lo
	131614	AB002438	Hs.29596	Homo sapiens mRNA from chromosome 5q21-2	lo-hi-lo
50	104933	N94126	Hs.12969	hypothetical protein	lo-hi-lo
50	302235	AL049987	Hs.166361	Homo sapiens mRNA; cDNA DKFZp564F112 (fr	lo-hi-lo
	320574	AL049443	Hs.161283	Homo sapiens mRNA; cDNA DKFZp586N2020 (f	lo-hi-lo
	324678	Al990739	Hs.77868	ORF	lo-hi-lo
	331022	H03109	Hs.108920	HT018 protein	io-hi-io
<i></i>	332430	H25350	Hs.21145	hypothetical protein FLJ22489	lo-hi-lo
55	330601	U90916	Hs.82845	Homo sapiens cDNA: FLJ21930 fis, clone H	lo-hi-lo
	101988	AF221521	Hs.8068	hematopoietic PBX-Interacting protein	lo-hi-lo
	102859	AL036058	"Hs.76807	major histocompatibility complex, class	lo-hi-lo
	101363	M11321	11 000000	1	lo-hi-io
60	133968	AA355986	Hs.232068	transcription factor 8 (represses interl	to-hi-lo to-hi-lo
OU	332530	M31669	Hs.1735	inhibin, beta B (activin AB beta polypep	
	317777	NM_014785	Hs.47313	KIAA0258 gene product	lo-hi-lo
	100452	D87742	Hs.241552	KIAA0268 protein	lo-hi-lo
	112988	NM_014867	Hs.5333	KIAA0711 gene product	lo-hi-lo
65	320848	AB020691	Hs.198232	KIAA0884 protein	lo-hi-lo
65	105162	AL133033	"Hs.4084	KIAA1025 protein	lo-hi-lo
	133905	AB028974	Hs.137476	KIAA1051 protein	lo-hi-lo
	331406	BE176893	Hs.23440	KIAA1105 protein	lo-hi-lo
	321441	AF107493	Hs.118498	Homo sapiens LUCA-15 protein mRNA, splic	lo-hi-lo
70	131913	AW207440	Hs.185973	degenerative spermatocyte (homolog Droso	lo-hi-lo
70	135424	U67611	N- 400704	transaldolase 1	lo-hi-lo
	128506	L40904	Hs.100724	peroxisome proliferative activated recep	lo-hi-lo
	330506	Al130740	Hs.6241	phosphoinositide-3-kinase, regulatory su	lo-hi-lo
	311251	AI655662	Hs.197698	ESTs	lo-hi-lo
75	314171	AI821895	Hs.193481	ESTs	lo-hi-lo
75	106096	AW379378	Hs.170121	protein tyrosine phosphatase, receptor t	lo-hi-lo
	133740	AW162919	"Hs.170160	RAB2, member RAS oncogene family-like	lo-hi-lo
	119521	W38038			lo-hi-lo
	119546	W38169		•	to-hi-lo
00	119559	W38197	11- 20070	estimableatoma hinding acatain 2	lo-hi-lo
80	133797	AL133921	Hs.76272	retinoblastoma-binding protein 2	lo-hi-lo
	305096	AA642964	Hs.163593	ribosomal protein L18a	lo-hi-lo lo-hi-lo
	120256	AA169801	Hs.98710	hypothetical protein	10-111-0

	200042		11 07/100	507	
	322919	AA178955	Hs.271439	ESTs	lo-hi-lo
	300566 330694	R34926 AI741617	Hs.326392 Hs.108447	son of sevenless (Drosophila) homolog 1 spinocerebellar ataxia 7 (olivopontocere	to-hi-to to-hi-to
	302416	AL120259	Hs.76691	stannin	lo-hi-lo
5	319289	AA037534	Hs.79059	transforming growth factor, bela recepto	lo-hi-lo
•	134656	AI750878	Hs.87409	thrombospondin 1	lo-hi-lo
	130117	U06641	Hs.150207	UDP glycosyltransferase 2 family, polype	lo-hi-lo
	124357	N22401		gb:yw37g07.s1 Morton Fetal Cochlea Homo	lo-hi-lo
10	108293	AA069155		gb:zm10f11.s1 Stratagene pancreas (93720	to-hi-to
10	108657	BE567753	Hs.132955	BCL2/adenovirus E1B 19kD-interacting pro	lo-hi-lo
	108658	AA641695		gb:nr62h10.s1 NCI_CGAP_Lym3 Homo saplens	lo-hi-lo
	331278	AA071383	th. 400000	gb:zm61d05.r1 Stratagene fibroblast (937	lo-hi-lo
	108340 108679	AA069820 AA115963	Hs.180909 Hs.323423	peroxiredoxin 1 ESTs, Moderately similar to B Chain B,	ko-hi-lo ko-hi-lo
15	108406	AA075424	Hs.325505	ESTs, Moderately similar to HBA_HUMAN HE	lo-hi-lo
10	114598	AA075601	110.020000	gb:zm88c05.r1 Stratagene ovarian cancer	lo-hi-lo
	108462	AA079347		gb:zm96c06.s1 Stratagene colon HT29 (937	lo-hi-lo
	108466	AA079409		gb:zm96h02.s1 Stratagene colon HT29 (937	lo-hi-lo
•	108489	AA082977		gb:zn07h10.r1 Stratagene hNT neuron (937	lo-hi-lo
20	330859	AA082977		gb:zn07h10.r1 Stratagene hNT neuron (937	ło-hi-lo
	108505	AA083376		gb:zn09g08.s1 Stratagene hNT neuron (937	lo-hi-lo
	331283	AA467736	Hs.275437	ESTs	lo-hi-lo
	100641	AW068302	"Hs.182183	Homo sapiens mRNA for caldesmon, 3' UTR	lo-hi-lo-hi
25	100642	AW068302	"Hs.182183	Homo sapiens mRNA for caldesmon, 3' UTR	lo-hi-lo-hi
25	325889 338038			CH.16_hs gij5867087 CH22_EM:AC005500.GENSCAN.149-9	lo-hi-lo-hi lo-hi-lo-hi
	338316			CH22_EM:AC005500.GENSCAN:145-5 CH22_EM:AC005500.GENSCAN:304-2	lo-hi-lo-hi
	100999	H38765	Hs.80706	diaphorase (NADH/NADPH) (cytochrome b-5	to-hi-lo-hi
	331131	R54797	115.50700	gb:yg87b07.s1 Soares infant brain 1NIB H	lo-hi-lo-hi
30	310955	Al476732	Hs.263912	ESTs	lo-hi-lo-hi
	311137	AW207582	Hs.196042	ESTs	lo-hi-lo-hi
	311598	AW023595	Hs.232048	ESTs	lo-hi-lo-hi
	313070	AJ422023	Hs.161338	ESTs	lo-hi-io-hi
35	110844	Al740792	Hs.167531	methylcrotonoyl-Coenzyme A carboxylase 2	lo-hi-lo-hi
33	120328	AA923278	Hs.290905	ESTs, Weakly similar to protease [H.sapi	lo-hi-lo-hi
	105914 129389	AW245680 NM_012445	Hs.9701 "Hs.288126	growth arrest and DNA-damage-inducible, spondin 2, extracellutar matrix protein	ło-hi-lo-hi lo-hi-lo-hi
	102759	NM_005100	Hs.788	A kinase (PRKA) anchor protein (gravin)	to-lo-hi
	100168	H73444	Hs.394	adrenomedullin	lo-lo-hi
40	102348	U37519	Hs.87539	aldehyde dehydrogenase 8	lo-lo-hi
	134158	U15174	Hs.79428	BCL2/adenovirus E1B 19kD-interacting pro	lo-lo-hi
	133908	AU076820	Hs.325474	caldesmon 1	lo-lo-hi
	101883	AU076743	Hs.75613	CD36 antigen (collagen type I receptor,	lo-lo-hi
45	327821		11 400000	CH.05_hs gl]5867968	io-lo-hi
43	134133	AA262294	Hs.180383	dual specificity phosphatase 6	lo-lo-hi
	103000 109251	NM_001975 AA194776	"Hs.146580 Hs.85935	enolase 2, (gamma, neuronal) EST	lo-lo-hi lo-lo-hi
	315566	AB037810	Hs.18760	KIAA1389 protein	to-lo-hi
	324697	AK000742	Hs.126774	L2DTL protein	lo-lo-hi
50	306011	AA896986		gb:al06a08.s1 Barstead spleen HPLRB2 Hom	lo-lo-hi
	307111	AJ174528		gb:an45g10.s1 Gessler Wilms tumor Homo s	lo-lo-hi
	106639	AV655272	Hs.20252	novel Ras family protein	lo-lo-hi
	106753	AI656166	Hs.7331	hypothetical protein FLJ22316	lo-lo-hi
55	107974	AW956103	Hs.61712	pyruvate dehydrogenase kinase, isoenzyme	lo-lo-hi
22	112033	R49031	Hs.22627	ESTs	lo-lo-hi
	113816 116024	H46008 AA088767	Hs.31518 *Hs.83883	ESTs transmembrane, prostate androgen induced	lo-lo-hi
	116158	AA381807	Hs.61762	hypoxia-inducible protein 2	lo-lo-hi lo-lo-hi
	119071	R31180	113.01702	gb:yh62b02.s1 Soares placenta Nb2HP Homo	lo-lo-hi
60	120132	W57554	Hs.125019	ESTs	lo-lo-hi
	120655	AA305599	Hs.238205	hypothetical protein PRO2013	ło-lo-hi
	122411	AW172356	Hs.99083	ESTs	lo-lo-hi
	320779	AA815354	Hs.169898	ESTs	lo-lo-hi
65	321024	AW246216	Hs.32058	Homo sapiens C1orf19 mRNA, partial cds	lo-lo-hi
65	321408	AW081530	Hs.137088	ESTs, Weakly similar to ALU1_HUMAN ALU S	lo-lo-hi
	323620 314946	AA306997 Al097229	Hs.268362 Hs.217484	ESTs, Weakly similar to hypothetical pro ESTs	lo-lo-hi <b>lo-lo-</b> hi
	320683	AA334511	Hs.26638	ESTs, Weakly similar to unnamed protein	to-lo-hi
	128959	Al580127	Hs.107381	hypothetical protein FLJ11200	lo-lo-hi
70	128896	T53925	Hs.107	fibrinogen-like 1	lo-lo-hi
	133592	AV652066	Hs.75113	general transcription factor IIIA	lo-lo-hi
	103245	BE566343	"Hs.28988	glutaredoxin (thioltransferase)	lo-lo-hi
	314785	Al538226	Hs.32976	guanine nucleotide binding protein 4	lo-lo-hi
75	103677	Z83806		gb:H.sapiens mRNA for axonemal dynein he	lo-lo-hi
75	131170	NM_014253	"Hs.23796	odz (odd Oz/ten-m, Drosophila) homolog 1	lo-lo-hi
	131164	AW013807	Hs.182265	keratin 19	lo-lo-hi
	100409 133167	D86957 AW162840	Hs.80712 Hs.6641	KIAA0202 protein kinesin family member 5C	lo-lo-hi lo-lo-hi
	319080	AW967646	Hs.23023	ESTs	lo-lo-hi
80	330706	AF097994	Hs.301528	L-kynurenine/alpha-aminoadipate aminotra	lo-lo-hi
- •	104052	NM_002407	Hs.97644	mammaglobin 2	lo-lo-hi
	100547	M57417		gb:Homo sapiens mucin (mucin) mRNA, part	lo-lo-hi

	103145	X66276	Hs.169849	myosin-binding protein C, slow-type	lo-lo-hi
	301015	AV655272	Hs.20252	novel Ras family protein	lo-lo-hi
	311013	AA224760	°Hs.153	ribosomal protein L7	lo-lo-hi
	132050	Al267615	Hs.38022	ESTs	lo-lo-hi
5	132349	AW975654	"Hs.181286	serine protease inhibitor, Kazal type 1	lo-lo-hi
	130889	AW972512	Hs.20985	sin3-associated polypeptide, 30kD	lo-lo-hi
	130791	AF030403	Hs.199263	Ste-20 related kinase	lo-lo-hi
	130385	AW067800	Hs.155223	stanniocalcin 2	lo-lo-hi
	127229	AA316181	Hs.61635	six transmembrane epithelial antigen of	lo-lo-hi
10	133820	S69681	°Hs.177582	surfactant, pulmonary-associated protein	lo-lo-hi
	129523	M13231	Hs.274509	T cell receptor gamma constant 2	lo-lo-hi
	321415	BE621807	Hs.3337	transmembrane 4 superfamily member 1	lo-lo-hi
	131859	AW960564	"Hs.3337	transmembrane 4 superfamily member 1	lo-lo-hi
	133444	M63978	Hs.73793	vascular endothelial growth factor	lo-to-hi
15	332567	AW939251	"Hs.25647	v-fos FBJ murine osteosarcoma viral onco	lo-lo-hi
	131328	AW939251	"Hs.25647	v-fos FBJ murine osteosarcoma viral onco	lo-lo-hi
	315901	AI521558	Hs.7331	hypothetical protein FLJ22316	lo-lo-hi
	104394	AA129551	Hs.172129	Homo sapiens cDNA: FLJ21409 fis, clone C	lo-lo-hi
	103739	AA115173		gb:zn30d02s1 Stratagene neuroepithelium	lo-lo-hi
20	103797	AA080912		gb:zn04d03.r1 Stratagene hNT neuron (937	lo-lo-hi
	103804	AA129196		gb:zn29d08.r1 Stratagene neuroepithelium	lo-lo-hi

## TABLE 18

5

Pkey: Unique Eos probeset identifier number CAT number: Gene cluster number Accession: Genbank accession numbers

5	Accession:	Gendank access	ion numbers
10	Pkey 108462 108489 101216	CAT Number 116651_1 118662_1 17379_1	Accessions AA079347 AA079506 AA079538 AA079442 AA082977 AA082955 AA082956 AA284166 AA314707 L25876 L27711 AA092745 N92087 U02681 AA315766 BE385121 AA352693 NM_005192 AI739135 A1066521 AW173105 AA257103 AA450169 AW261971 AA305065 AI954494 AW950384 AW732122 AA830348 AA789097 AA777794 AA284072 BE564465 AI005313 AA804528 AI041134 AI700317 AI352491 AA856987 AA769007 AA494334 AA769862 AA831168 AI143496 BE090796 AA831166 AI141222 AI372907 N64843 AI075136 AI076701 AA464156 AI076409 AI273523 AA627383 BE043332 T95666 AA158102 AA158059 AW340182
15	131328	8509_1	AA257019 Al206700 Al678081 AA757304 AA055005 AW059834 AL039012 AW939251 NM, 005252 AU076596 V01512 V01512 AW579056 AA249247 Al590359 AW510478 AW518282 BE046054 AW874080 Al268596 AA996237 Al695592 Al244117 AA290764 AA401957 AA50878 AA428304 W74018 W74016 AA040944 Al272071 AA745909 AA620979 AA019816 Al245094 AW009706 AA662536 AW024264 Al268601 AA932024 AW513222 AW024169 Al659705 AA932526 AA975329 Al567603 Al889320 AA514238 AA020837 Al623966 AA843677 AA477453 AA496353 AW372625 AV656426 K00650 W96348 N62388 R95977 AA434270
20			A1093633 T27639 AW960245 AW881177 R15253 N36936 F07701 AA319315 AA337290 AA284642 AA344052 F05184 AA351062 AA378451 AW794233 AW884380 N36951 R49879 AB022276 AA300350 AW839435 AW191708 BE220350 AA280404 AA485546 AW794235 AV654223 AW838891 AA285986 N752823 AA335648 AA371089 AW845414 H63166 R12840 AA379680 AA477579 R13148 H71003 H71015 AA362156 AW750674 AW845415 AA366924 AW608044 AI570388 R31511 R33906 R33921 AW663022 AW360985 AI207838 AW607239 AI672451 AI573282 AW794752 AA370328 AW998896 AW797239 AW998912 AW794742 AI955434 AI810067 AW073373 AA370325 AW195330 C18106
25			AW998736 R79476 AA429721 Al891081 Al381534 AW022137 AW020000 Al630329 N99428 Al870222 Al971257 Al922196 Al857753 AW579397 D56749 Al925005 Al685727 AW805573 Al982678 Al784604 Al005625 AW877772 Al634947 Al950829 AA493243 BE166086 Al801820 Al925643 Al627992 AW316704 Al261318 D57757 AA887178 AW770406 Al972075 Al222254 Al675794 D58060 Al701954 D58166 Al799500 AW805669 AW276098 AW874253 Al962991 Al248184 AW996924 Al017462 AW022260 Al885957 BE176841 AA878863 Al697419
30			AW662094 Al479529 BE177025 D57403 AA507952 AW664593 AW800998 AI985773 AA566089 AA442759 AI624670 AI460284 AI800205 AI537788 AI537593 AI244382 AA583463 AA922678 AA864382 AI610837 D58070 AA844283 AA947992 N73801 AI453821 D58184 AI678887 AW243755 AA746085 D57742 AA757380 R44148 AA496403 BE180303 AW363556 BE006616 D57395 AW805507 AW805511 AA617991 AI373585 H30122 D57744 AW805501 D57691 D58148 AW873164 AW768483 D57601 D58795 AA837997 BE180123 D57599 AA485387 AW022208 D58096 N67917 W95944 AW805506 D57518 D57990 AI074096 D56521 D58151 AA428720 D56648 D57778 AW805504 D57750 D58108 AW021706 D57449 D57041 D58277 D56935 AI356974 D57023 AA018712 H27631 D57851 D57514 D57268 D57468 AW805646
35			A278945 D57323 D56986 D57639 D57829 D58078 AW805515 Al348684 D57772 R74449 BE041558 D56746 AW805869 A278945 D57052 D56849 D56840 A278945 D56746 AW798485 D56640 A278945 D56702 D56849 D56847 AW581419 AA470397 D57591 AW798984 T27640 N66497 D56803 AA618186 AW805647 D57945 N23726 D56637 N23730 D56992 BE176889 BE176899 BE176909 D56757 N68137 D56987 Al559806 AA631437 D57464 D56718 C17030 T29278 D57377 AW021936 AW118330 AA515358 D56610 AA494092 D56934 T97774 AI473546 R74350 R84834 AA579200 D56616 C03207 D57391 N52416 D56928 R79209 D56925 AA020879 D45546 AI858769 R20750 T09381 F01435 AW627906 D58202 Al933993 F01912 H27552 AA174191
40	124148	31218_1	T16515 AW023216 AA434146 H83387 Al346751 V01512 V01512 AA576407 AW365140 AA937471 BE174681 Al568829 Al274663 R85530 AL048225 H83388 AW798734 BE300994 BE384439 AW794648 NM_002305 M57678 Al929016 AU076727 Z83844 Z83844 Al906100 W44519 H98497 AA188069 AA572687 AA035793 W93978 BE409220 AA359751 AA502475 H28319 AA527889 AA432335 AA864762 AA340061 C05180 W68192 AA327811
45			AA345871 AI750205 N34093 N86639 AA085753 AA603415 AI355561 AA442262 N42135 C04367 N57266 AI038364 AI184846 AI928853 X15256 J04456 AA603552 AA317300 AA588615 AA813495 N40276 AA400624 AW264988 H21418 AA643822 AA603569 AA507955 N44497 AI000869 AW079049 AA614629 AA303987 AA362817 H54502 N85495 W52256 F30576 AA568129 H26935 W93977 AA373651 AA872398 AI332540 AW572787 F20782 AA442263 AW301076 AA558556 AA825366 W23842 AI038829 AA302408 AA374629 AA614477 AA341686 AA374846 AA187091 F24764 AA157099 AA374853 AA991592 F26839 AA744090 AA936881 AA374627 AA329755 AA854398 AA618108
50			AA973600 AA757956 W44520 AA379779 AA373698 AA369135 AA380039 BE408327 AA375117 AA375744 AA380014 AA373556 AI335987 AA903267 AA828223 F25088 AI246573 AA299386 BE275844 BE275666 BE384214 BE620707 AA975886 AA858048 BE548468 AA193055 BE274324 AI870164 AA129614 AA922761 AA935745 AA374567 AI580916 AA3745661 AW239224 AA374466 N52172 F24306 AA30453 AA363443 AA5868627 F19159 AA580021 N90877 AA654335 AA679168 AA573071 AW238834 AA988739 AW239423 AA976330 AI074239 AA999911 AI200930 AI971173 AI187321 AA937760 AI016242 AA373684 AI094874 AI302174 AA641237 AI370974 AI971010 AA400379
55			AA679137 Al096579 Al001918 AA524101 X14829 AA081302 N30374 Al338782 W74444 AA528232 Al734954 AW188024 AA433857 W92348 W94431 Al708356 Al753458 AA494460 AA825257 AA614246 AA039477 Al350213 Al309110 AA745965 AA291936 AW001376 Al066764 W74407 F30627 AA291937 AA480615 AA931667 AA331315 Al936154 AA824332 AA181109 Al017291 AA934736 AA062637 AA599977 H54814 AA635624 Al802655 AA564078 R69997 AA716551 F30469 AA961030 Al126757 Al183943 Al066798 Al419436 AA302095 AA157768 AA953030 AA588476 AA131216 T79619 Al752885 AA614820 AA988962 Al143561 AA493182 Al302481 AA301613 R73520 AA069898
60			AA374944 AW364221 AA342013 Al244949 F36390 AW050980 N79486 AA101160 T68112 Al750204 AA328787 H02617 AA314734 AA527923 AA307835 Al885112 Al872905 AA534666 AA188363 Al192490 H45772 Al824700 Al184276 AW079473 N29847 AA720843 AA720914 AA573391 H54416 T59424 Al824457 AA304220 AA482553 W72882 AA627932 H27514 H28400 W68050 H290953 AA635786 H21376 AA514046 Al342823 F29905 H25999 AA757144 H21636 F22104 AA428650 F27143 F283346 AA535690 H45771 AA548851 AW170154 H45646 W92274 Al921614 AA176461 AW170153 Al927284 Al161206 AA594439 T28595 H41129 Al497579 AA978015 AA328875 AA373653 AA090973
65			AA328623 AA328759 AA366468 AA375406 H46976 R86050 H02722 AA328321 AA328205 R62358 AA373717 AA304138 AA304224 AA301603 H54867 AA374783 AA376232 AA373239 AA374917 AA375573 AA303857 AA376466 AA302613 AA302613 AA304082 AA301731 AA357988 AA303328 R25744 AA301587 N78746 H20508 AA659423 R47960 AA825456 A1001806 A1245114 AA729223 AA860271 A1913845 H26296 AA733035 AA340965 AA304291 H27356 H20598 AA129613 R69996 AA157689 H20992 W16630 W16561 H25964 H21754 W01159 W42885 AA176730 H39504 N39788 AA182956 H27585 AA082164 AA328927 AA339934 H61805 H61804 H45580 AA476229 AA714104 AA507471
70			Al262184 Al139474 Al139476 Al001045 AA614374 AA593153 F33347 F34679 T68225 N25703 AA186999 Al623318 F18313 N72069 AA903161 H38546 H28672 Al880529 Al128960 AA299183 AW768886 F17445 F30433 AA303984 AA303687 AA309366 H28320 Al659479 AA627222 AA064882 AA507447 R53171 AA039476 T79704 R36589 T83222 H26453 AA298798 R53415 N84918 F37846 R94423 AA352679 AA308615 AA375442 BE173864 AA353674 R73519 R62478 T59480 AA089852 Al265789 Al077675 T90770 R54006 H46977 AA187168 AA157123 H21637 R48072 AA814207 R53082 AA305829 R62359 Al818429 AA887755 AA534238 Al813821 AW023928 AA062712 Al698995 F19074 AA345870 Al658776 AA903325 S44881 AA379844 N86780 AW089895 F29687 W52257 AA131229 AA978007 AW953024 R94945 H28332
75	124153	25750_1	AU077333 M81635 NM_004099 X60067 Al686183 AW401439 T39535 AA302410 AV645727 AV653397 AA317395 AA218582 AA219682 AA227317 AI750900 BE440055 H77491 F12371 AA314714 T74055 Al655647 AA489421 AA346569 AI129523 AA094975 AW793582 R97358 H67966 N72440 H79590 H81459 H60508 R39523 H60900 H40547 AA377244 AA318430 H71201 R64651 R65629 H72546 AW798947 N76974 H03029 N77701 AW151751 H60925 AA455839 H72947 N58334 N55487 AI29891 AA581634 AV651323 AV651728 AV650086 AV651295 AV648042 AW020600 Al537887 AA429713 AW080244 N73463 AA471335 AW150316 AA360851 W01407 BE074301 W21371 T87221
			AA190691 D16906 AW862400 AV661466 A357816 AA442743 A1189966 AW887793 BE005206 AJ926016 AA317024 AA976151 AA247314 A767184 R64644 R62817 D57965 N74437 N74385 H60409 N66059 H91165 R79462 F09991 R26175 H77853 N32590 D55667 AA461122 D56666 D56903 AW021856 AA374084 R69734 H66894 T81638 T63958 W23935 R67668 AW021682 H81249 H61959 H89852 R79306 W25710
			109

W42964 AA384428 AW994316 H95163 H95158 R33688 W46557 AW748451 AA029916 AA463826 AA314287 R23084 AA368891 H02926 AA310456 H03632 C02397 R63745 H94539 R32226 R24648 H44502 AA039571 AA345336 W42846 R48024 R79724 R63143 AA379513 R21780 R80704 T70422 H21580 H46388 R62779 AA579734 N64111 AA344527 Al865473 R66666 Z20058 T52284 H95103 R36513 R21874 R31363 AA220939 BE439695 A189683 AA164901 Al539383 AA768249 AA442361 W02867 AA303315 AW952009 AA314544 Al076799 AA216780 T70338 AA039672 AW629489 AL044620 AA533203 AA043082 Al668619 AW298204 AW195268 AI391606 AA437282 AW304801 AW085720 W02586 AA863279 T82339 AI356879 BE464557 AI038992 A190018 BE146083 Al860399 AI039572 AW26873 AW468134 ANA02874 AA042874 AA042874 ARA02874 AA042875 AA032874 AA042875 AA032875 AA032874 AA042875 AA032875 AA032874 AA042875 AA032875 AA0328 5 AW06972 W3505 Ax65239 Ax6527 16239 A35677 A1754185 AA680298 AA460262 H9132T7 N57879 R66069 N95684 AA040855 AA227116 N94486 H04229 H97877 A1161080 A1074367 A1025767 A1754185 AA888150 A1356979 R79463 AA029917 R69637 A1810134 AA460820 A1377990 A1743170 AA854637 AA628548 AA664223 A1362196 AA489363 A1361404 A1363155 AA300504 A1678269 AA633851 H61743 A1161012 AW339721 W42847 W46558 AA143120 A1042475 AA479365 AA219592 AW468142 H67690 A1186516 AA531387 AA835378 H03030 T68119 10 AW339721 W42847 W46558 AA143120 Al042475 AA479365 AA219592 AW468142 H67690 Al186516 AA531387 AA835378 H03030 T68119 H95133 AL040491 Al289149 R63701 R32177 R32865 Al811374 Al613274 AA775300 AW192882 R37509 W42965 R47918 Al949525 Al129450 H49378 Al435907 Al832271 AA478271 R21849 H03633 Al888539 C75673 Al261334 AA614478 AW469307 Al261429 W03148 AW026141 AW236371 R79725 AA346568 C06197 T27764 H59538 Al749196 AA485299 AA719227 Al698762 N70790 Al925028 R21734 AA977432 H77905 Al625648 AA918868 Al220069 Al352568 AA668729 AA195395 T63334 Al932783 N32271 R26048 H90697 R24539 Al970287 T55374 N93019 T11162 AA377400 AW882126 AA602293 F35923 Al424237 Al826517 H27442 AA039729 AA382630 Al567304 AA045112 T57779 Al474576 Al352569 R63095 H44456 X85116 Al521609 AA164352 BE146079 H60082 Al334776 AA700506 AA782742 R67386 R22978 R33584 R67011 R80705 Al245311 H81590 Al360786 Al219244 R39564 H66850 Al184385 AA687691 H68013 AA092081 Al4445480 AW005734 AW068302 Al754558 Al750727 Al752631 AA30174 AA327522 M64110 AW859944 AW859989 Al751995 AA769620 Al858829 Al924875 Al888836 AA684291 Al685060 AW088029 Al924908 AW466328 Al093800 AA991651 Al254501 BE004703 AA334442 AW938852 AA194330 Al046853 AA857866 AW391995 W30846 AW66329 AW266328 Al093807 H90607 H00703 AA354444 AW938852 AA194330 Al046853 AA857866 AW391995 W30846 AW66329 AW266328 Al093807 H90607 H00703 AA354444 AW938852 AA194330 Al046853 AA857866 AW391995 W30846 AW663298 W25761 AA042863 R90965 H97060 W03910 H94687 T88984 AL048165 T29632 N31556 15 28620\_1 100641 AL046953 AA852866 AW391995 W30846 AW662928 W25261 AA042863 R99045 H97060 W03910 H94687 T88984 AL048165 T29632 N31556 20 N36484 A1798679 AA989355 W23832 AA873789 A1743646 AA363587 A1814748 AW338990 N73740 N83666 AL047816 R24137 R63433 NSSSSS VANDER NR NSSSS VANDER NR NSSSSS VANDER NR NSSSS VANDER NR NSSS VANDER NR NSS VANDER NR NS VANDER NR NSS VANDER NR NS VANDER NR NS VANDER NR NS VANDER AA527960 AA525036 AA044414 AI752460 AA703064 R01216 AA897183 AI751996 T81078 H95047 AA573642 D58348 N20953 AA437143 N95439 AA579540 AW867056 AA770090 Al085180 Al806799 AA426421 Al572513 R24081 AA853189 AA295620 AA234044 AA371020 AW994984 H20896 AW964438 AA318516 AA318499 AA318727 AA318211 AA318478 AA318444 AA318307 AA318497 AA318448 AA318309 AA318496 AA318243 AA318435 AA318424 AA318217 AA318523 AA318438 AA318487 AA583185 AW994985 T69842 Al251813 AA478174 AA447737 T68350 F07712 AA121145 H08973 AA345212 BE000667 AW068210 AW608407 R05674 H16712 N85426 N42354 H85516 BE147991 T28113 R32662 AA384678 AW239275 H82382 AW840700 D58229 C04082 W45394 AW795667 R73973 BE0022409 AA042828 AA365355 AL223812 AA344709 BE149590 R70995 W46881 W90778 N71242 AA534826 AL040676 R23797 H96450 AA062957 D79947 W46960 AW959278 AA295997 AA026215 AW579469 AW365135 AW365134 AW994353 AW972886 AW069166 AA343690 AW888731 AI751527 AA937490 AA937506 Al826715 BE465604 Al925532 Al858109 AW339097 Al858524 AI720571 BE046506 AW384981 AA043908 AA375983 AA525181 AW068366 AW070577 AW891837 N83985 AW182753 Al422979 AI679733 BE006555 AL048166 Al081401 Al888821 AI626043 N37087 AI624140 Al801298 AA600048 AI753947 H89615 N66424 AW069796 AI814880 AJ982806 AI754287 AI971816 AW0690022 AW069069 AW0669454 AA342989 AI077712 AI311467 AI087361 AI801015 W46993 AI281324 AW191963 AI421675 AI300881 AI356670 N95439 AA579540 AW867056 AA770090 AI085180 AI806799 AA426421 AI572513 R24081 AA853189 AA295620 AA234044 AA371020 25 30 AW069069 AW069454 AA342989 AJ077712 AJ311467 AJ087361 AJ801015 W46993 AJ281324 AW191963 AJ421675 AJ300881 AJ356670 35 AW069069 AW069454 AA342989 AI077712 AI311467 AI087361 AI801015 W46993 AI281324 AW191963 AI421976 AI300881 AI359670
AA873156 AI004219 AI189685 AA478018 AA076063 AI445222 AI753124 AI521569 AI925026 AI022368 AI475993 H20846 AI222324 AI635123
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AI370492 C16471 AA652809 AA936687 AA506512 C16306 AW028413 AI537935 AA528347 C16255 AW029046 C16202 AI868152 AI524662
T94414 AI567041 AI619654 AW008486 AI075624 AA577434 AA693617 AA979107 AA95774 AA85045 AW773763 R45484 AI570898 L154708 T49285 40 1941 A AISCHOP AND 1964 AWU05469 AU 5824 AV 5745 AND 1954 45 AV559047 AV659632 AJ750389 AA092053 AA092798 HB5367 T61597 R23745 Z20418 T78485 AJ751528 AW068121 AA853188 AJ752459 AA853711 AW950663 R78964 R36359 R21626 R21522 AA853711 AW950663 R78954 R36359 R21626 R21522

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AI888836 AA864291 AI685060 AW088029 AI924908 AW466328 AI093800 AA991651 AI254501 BE004703 AA334442 AW938852 AA194330

AL046953 AA852866 AW391995 W30846 AW662928 W25261 AA042863 R99045 H97060 W03910 H94687 T88984 AL048165 T29632 N31556

N36484 AI798679 AA989355 W23832 AA873789 AI743646 AA363587 AI814748 AW338990 N73740 N83666 AL047816 R24137 R63433

AA524984 AA234043 AA195131 N99903 AA453669 AI240302 AA370271 AI950026 AW771049 AA121476 AA569557 AI752632 AI355594

AI471993 AI159941 N94555 AI753138 N21537 H97881 N25769 AW068044 AA808425 R63380 AA384736 AA384738 AA852352 AI073645

AA527960 AA525036 AA044414 AI752460 AA703064 R01216 AA897183 AI751996 T81078 H95047 AA573642 D58348 N20953 AA437143

N95439 AA579540 AW867056 AA770090 AI085180 AI806799 AA426421 AI572513 R24081 AA853189 AA295620 AA234044 AA371049

AM804094 N20986 AW867056 AA770090 AI085180 AI806799 AA426421 AI572513 R24081 AA853189 AA295620 AA234044 AA318409

AW804094 N20986 AWR647056 AA770090 AI085180 AI806799 AA4384718 AA318478 AA318408 AA318479 AA318448 AA318497 AA318448 AA318497 AA318448 AA318479 AA318448 AA318479 AA318448 AA318479 AA318448 AA318479 AA318448 AA318479 AA318444 AA318309 100642 28620\_1 50 55 AW994984 H20896 AW964438 AA318516 AA318499 AA318727 AA318211 AA318478 AA318444 AA318307 AA318497 AA318448 AA318309 AA318496 AA318213 AA318435 AA318424 AA318217 AA318523 AA318438 AA318487 AA318724 AA593185 AW994985 T69842 Al251813 AA478174 AA447737 T68350 F07712 AA121145 H08973 AA345212 BE000667 AW068210 AW608407 R05674 H16712 N85426 N42354 H85516 BE147991 T28113 R32662 AA384678 AW239275 H82382 AW840700 D58229 C04082 W45394 AW795667 R73973 BE002409 H85316 BE14/991 128113 K32662 AA384678 AW2392/6 H8238Z AW844//UU D36229 C0408Z W45394 AW795667 K73973 BEUU2409
AA042828 AA363555 AJ223812 AA344709 BE149590 R70995 W46881 W90778 N71242 AA534826 AL040676 R23797 H96450 AA0622957
D79947 W46960 AW959278 AA295997 AA026215 AW579469 AW365135 AW365134 AW994353 AW972886 AW069166 AA343690 AW888731
A751527 AA937490 AA937506 Al826715 BE465604 Al925532 Al868109 AW339097 Al858524 A1720571 BE046506 AW384981 AA043908
AA375983 AA525181 AW068366 AW070577 AW891837 N83985 AW182753 Al422979 Al679733 BE006555 AL048166 Al081401 Al888821
Al626043 N37087 Al624140 Al801298 AA600048 Al753947 H89615 N66424 AW069756 Al814880 Al982206 Al754287 Al971816 AW069022
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	` 130791	30310_1	AF030403 AF099989 NM_013233 AW104402 AA251775 AA251558 AI582744 AI222132 AI351849 AA150838 AA905073 AA278308 AI830043 AI803232 AI813651 AI858774 AI266366 AA286879 AA587082 AI351439 AI080241 AI873470 AI276052 AI392761 AI018158 AW195899 AW274293 AW592760 AA913004 AI936691 AA766905 AA648820 AA824515 AI016857 AA815184 AA642482 AA150717 AA332969 AA286878
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			Al948838 AW235336 AW172827 AA095289 BE046383 Al734240 W16699 Al660329 Al289433 AA933778 AW469242 AA468838 AA806983
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65	109220 116448	139161 <b>_1</b> 30623 <b>_</b> 1	AW337859 AA459672 AA729743 F02479 AW455759 AA729543 T25454 H62547 N50430 T63976 N70049 N54292 T63965 R85599 N49681 D17232 AA910951 W24824 BE386617 AI525551 BE567611 AL034410 Al198816 W73728 AA534300 W61040 AA102496 AA771826 N92556 H50961 N44829 AA628033 AA642158 AA628038 AI613134 AI468860 N87245 R91859 AA091252 AW958181 AA196018 AA706922 AA775199 AI798729 AA192334 AA132242 AA132243 AA973154 AA376742 BE268321 BE270484 BE268500 BE410641 AW247710 AL160131 BE304734 BE559730 BE385420 BE296695 BE270916 BE560389 BE513878
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75	10000	.002.	BE044507 Al936899 AA746205 AW005627 Al245057 Al651847 AW197857 Al420928 Al381889 Al283588 Al219782 Al355977 AW166497 Al090179 Al219676 Al681608 Al830080 Al418165 AW044649 AW294136 AW298132 AW292753 Al186178 BE326292 Al927366 AW589975 Al698895 BE041614 AW008628 Al634520 Al420668 Al272881 Al300751 Al914076 AW149564 AW663108 AW080009 Al336521 Al494509 AW117602 Al651038 Al770019 Al417431 BE222987 Al083931 Al672740 AW055034 Al826705 AW468862 Al873323 AA128147 AA128116
80	108647	10525_1	D57622 AJ307385 AJ559899 AJ537834 AA12688 F09994 AA026940 N56214 AA564903 D20488 H87549 AA868491 BE546947 NM_017409 AF255675 AA299577 AA314165 AW961165 AA307551 AW961168 AA659084 AJ673757 AJ796361 AJ670876 AA190344 N42572 BE076253 AA910251 BE621922 AJ796528 AJ458102 AA502954 AW024150 AJ653810 AA411006 AJ743397 AA190345 AA888101 AJ174335 AA916542 AA112396 AJ307395 N31842 AW205660 AJ269376 AJ129087 AW080195 AW024474 AJ369480 AW769611 AJ382520 AJ942373 AW469953 AJ949161 AA865803 AW994250
	108657	23226_7	BE567753 AA084916 AA113136

	108658 115881	112832_1 29310_1	AA641695 AA113139 AA074156 AA083045 AA074392 AA083158 AA113057 AA084807 NM_005756 XB1892 AA397668 W32664 AA436725 Al452634 Al003488 Al521155 AW274256 AA634329 W32478 AA435577 Al908762 Al289997 AA782155 AA730762 AA730771 AL045809
5	102012	21793_1	BE259035 AU077338 U03057 NM_003088 AW732835 AL134784 AL120159 BE409858 T09062 BE297271 BE294908 BE273148 BE259718 BE261678 BE261678 BE260498 BE408153 BE259762 BE261974 BE260884 U09873 AW410121 BE019189 BE278692 BE252072 BE383265 BE263157 BE262507 AA774906 BE296630 AW379502 BE538093 AA040533 BE297056 BE293964 BE297011 AA428510 T27582 BE262958 BE382541 AA077541 R87539 AW905865 AW905869 T49230 BE272643 BE256870 BE279288 AI940330 AI940368 BE019144 BE298451 W47256 BE272651 BE018948 BE255797 BE393014 AA852153 BE312227 BE262095 W07585 AA043912 AA403111 AL134704 AA459745 AA027019
10			M78875 BE265292 AW168964 AA451618 AA186594 AI187107 AI885355 AI339462 AI090054 AA040756 AI937569 AW055162 AI336276 BE205855 AI887647 W30953 AI375605 W95365 BE207928 W32489 AI887443 AA040292 AI093103 AI924761 AA437003 AI564843 AI369291 BE300843 AI566221 AI500381 BE312463 AI453117 AI885669 W78133 AA677779 AI2141855 BE208344 AI627739 AA450215 AI150235 BE206671 AI475936 AI493672 AI031615 AW467734 AA908886 AW297260 AV300302 AW515364 BE207810 AA461327 W68528 AI401541 AW298376 AI223266 AA157956 AA728777 AW268529 W81199 W70220 AI359697 AW360928 AI362260 AI961854 AI453087 AI922508
15			Al749454 AW771707 AW452557 W70219 R62745 AA041210 W68814 AA627393 Al218944 Al887035 Al088922 Al550788 T48210 AA564373 Al208904 Al950808 AW467727 Al081938 Al743346 Al016931 AA502297 Al924504 Al568845 Al671213 AA554629 Al650618 AW015272 Al283991 Al568493 Al968376 AW594745 Al341863 AW196605 Al65605 AW006227 Al087087 AW439650 Al085505 Al656126 Al001102 AW410122 AA927034 Al814950 AW470573 Al568575 AW073874 AW196325 AA665476 Al654701 Al364353 Al458249 AW150618 Al864978 AW103699 Al500694 T06432 AW304384 AW025677 Al682728 Al928669 AW072118 W95376 AA931596 T49231 W47384 M62123 Al880115 Al695915 BE551908 AA599588 AA587443 AA627613 Al825423 Al434053 Al961070 Al341081 BE019156 Al928667 AA923443 Al950232
20			AIG53915 BEG51906 MA393986 MA361445 MA621613 M625425 AIG54055 MISG1070 M3341061 BEG19136 MISC6007 MA323443 MISG6232 AIG53812 AI990944 AW337670 AI933222 AI985976 AA903555 AI719748 AIG21100 W02798 W02155 AA078409 AA922965 AI146961
	132116	96515_1	AA838643 AW044594 BE241997 BE258694 BE311788 BE259835 BE313211 AW960474 AA328243 AA704789 Al088169 N20591 Al823476 H81760 AA406184 W44795 Al040999 AA035348 AA632324 BE295273 AA815436 AA406294 AW394165 AA094618 BE296595 Al659092 AW297091 AA401881 Al435984 R80433 Al948677 W05276 AA234767
25	102034	598_1	H87282 AA253183 AI903474 AI903475 X72913 AL036029 AI903264 AI903383 AI903473 AI903426 AI903263 AI903331 AI903348 AA348154 AA558044 X75546
23	102034	390_1	NIM_002023 AA018495 AA568437 AA336463 AA336865 U05291 AA338073 AA360007 H26478 AA151040 R54231 H06222 AA411610 BE184970 R08802 H14537 BE184886 BE184857 BE184972 BE184861 BE184884 BE184890 AA283616 AA486471 H14444 H45124 H42970 H28253 H25914 AA009480 BE184935 AA305772 BE184889 BE184991 H28266 T27990 BE184880 AW950249 BE184965 BE395547 W65326 T49139 AA194473 AA194461 AA129907 AI680740 AW513000 AI346045 AA581716 AI674688 AI923173 AA587387 AA411190
30			AW474604 H06223 AW339985 AA908830 AI143335 AI806156 AW073728 AI570719 AI806149 AI653183 AI138299 AI807167 AI826341 H42900 AI925436 AW190948 AA150949 AI342245 AI991294 AI961468 AI360927 AI264267 AI015857 AI493989 AA527366 AI127268 AI304378 AI911417 AA682520 AI340130 AI346002 AI446304 AA916776 AA621369 AA129908 AI693879 AW836294 H26317 AA469443 H28206 AI609744 AI347112 BE218476 H45427 AA971098 H24697 AA861539 AA947902 AI339991 AI015777 AI828473 AI298202 W51828 AI298200
35			AW148725 H28218 AI440179 AA018496 AI249821 AI801917 AA663157 AW513059 AI827878 W61309 W48716 AA485748 T49140 AI168001 R08803 AI061052 AI804537 R51837 AI127238 AA284975 AA722722 AI520874 AA993797 AI039374 AA776215 AI084307 AW276143 H44534 AI214418 AW002315 AI422575 AI393603 AW449955 AA029408 AI582937 AI457747 AA194376 AI217628 AI432125 AI474369 AA911062 AI022596 AW276107 AI708968 AA628771 AW263915 AW150018 AA723100 H22132 AW051115 AI424515 AA526379 AI581791 AI933696 AI916839 AW003461 BE502206 AW593940 N67534 AI473431 AA501983
40	102076	28044_1	BE299197 Z85996 AW247234 AW249122 U09579 AW245698 AW250360 AW250483 BE241887 BE244900 AW247093 AW247357 AA380910 BE208575 AW583068 BE018355 AI751660 AA853842 BE206983 BE207145 L26165 L25610 U03106 S67388 BE257775 AA456445 W01311 BE263645 BE279085 AW732606 BE263622 BE265001 BE297240 BE263520 BE279288 BE256088 BE255900 BE252329 AW239199 L47232 L47233 BE207178 T08399 BE252557 H24262 BE258576 BE251231 AA381909 AW836368 BE206752 AA029109 AW820448 AW820447 AW843746 AA376396 AA321901 AA310434 AW249019 R46847 AW843743 AA375906 D31116 AA376199 AA065009 AW842884 AW842793
45			Al752795 W02824 H83378 M79002 R74301 BE545783 BE266009 R87600 BE180828 AA375519 AA376322 T53381 W39472 W44813 AA902104 R79427 AW246239 N44912 AA376207 R27374 BE253553 BE207052 AA376096 AW674390 AA187864 AA382005 N42395 N43766 AA373406 AW167163 AA320920 AA187865 BE245429 AA725216 T73450 AW248657 AW248580 AA614342 AW248690 AA134592 N95402 C06087 AW890404 AW890491 W74355 AW890497 AA481474 BE183394 AA854012 AA574254 AA84586 AW795988 AA029195 AA618214 W24029 N23941 AI753303 N35012 A1094940 W76550 N35823 R27375 AI123889 N33064 AI590965 AI146429 AI818625 N33420 D25587
50			AA716735 N34995 N35249 C21315 AW996039 AA302439 AA134593 AI494143 BE504194 AI191867 AI018663 AA554359 AI041908 AI369918 AA320526 AI041930 AA631887 AA629728 AI752794 AA857879 AI090388 AI751659 AA932518 AA705773 R46753 W45365 AW192512 AA603001 AA069286 AI355869 AI986250 AI813372 W45293 BE301229 AW511449 AW338097 AW247845 AA454553 AW246700 AA621693 AW780118 BE301970 AW129949 AW001364 AW732086 R79428 AI640886 AI814746 AI671072 AW338463 AI457683 AI955722 AW250453 BE046057 BE301255 AI623838 AI066427 AA069702 AI754612 AI274183 AI818343 AA838666 AI087209 AI859858 AI590130 W46516
55			AI565171 AI862747 AI453163 AI815083 AI628269 AI073360 AI572101 C05955 BE501618 AI677993 AI354988 AW337646 AI623382 AI890317 AI678661 AI979004 AI634583 AI867024 AI961540 AI973138 AI215141 AI866715 A862447 AW073998 AI365137 AW798270 AI813378 H22870 AI500446 W46448 AW591391 AA083831 T29792 AI952615 AI086902 T53382 R87514 A1147453 AA972341 AW839649 AI971521 BE257943 AI921309 AI079598 AI808947 AI872642 AI270535 AI826937 AI460140 AW615599 AA725669 AW250749 AA324110 AA310402 AA070728 W80687 AI283834 AW674560 AW572555 AI590414 AA534613 AW615591 AI274741 AA576395 AI148273 AI473483 AA644199 AA069979
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	132160	128565_1	W26406 AW136872 BE349103 AA935418 R54810 AI804000 AA879147 AI912294 AI339626 N40443 AA807907 AA446520 AA418512 AA101321 AA281770 AA227954 AI435989 AW975199 AA253044 R42784 R44804 AA227789 AA253099 AA280126 AI383274
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75			AW630441 AA115019 C75681 Al754224 AA164967 Al439457 H89682 AA983296 AW662585 AA853277 AA333773 Al754603 W30982 AA853377 BE169188 BE169223 AA115490 AW385682 AA993543 C02404 BE169185 BE169226 AW023640 R64182 AW993827 Al376301 AA043545 H62370 AW069486 D16887 AA343887 AA343832 AW9565555 BE175910 N47487 R69794 AW592387 T40161 AA884315 N98463 Al051407 Al678677 Al927978 AW664579 Al590727 Al090077 Al061451 Al147531 Al620584 AA617746 H99816 N47488 Al823867 AA605292 N64131 Al948780 Al767933 Al559598 Al564677 Al269461 Al420317 Al831711 Al146359 AW168879 Al699354 Al346276 Al818252 AW627427
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			110

			BE122814 AW630254 AW403442 R29627 AA375795 AW401663 AA683275 AA356222 BE410618 AA303830 AW498613 AW402378 BE258279 BE440060 BE539170 BE547722 BE620522 AW239013 BE618205 BE622304 AA381037 BE537289 BE538975 AW238805 BE546710 AA330972 AA376336 BE513559 BE295994 BE544569 BE543720 AA317908 N88057 BE545225 BE293935 AA360992 BE408886 AA206192
5			BE206839 BE398082 N84521 BE547881 AW651808 BE315212 BE545550 BE277333 BE274843 AW239248 AA176376 BE542891 AA348058 BE544201 BE548621 BE304755 BE269329 BE548052 BE269921 AL048472 BE560089 BE315560 BE613006 AW401848 BE293081 AA223405 AA356051 AA315114 BE267163 AW610536 BE409389 BE513915 BE270766 BE561005 BE560681 BE548838 BE546443 AA383753 M78064 AA376402 AW988176 A133342 BE614058 X03077 AA885733 BE548461 AW663131 AA853435 AA324697 AW405517 AW374649 BE019816 AW403098 BE539709 AW673496 AA373182 AA376953 AA090902 AW818624 AW393154 AW363737 AA143265 AA522902 AA375189
10			AW631262 N85647 BE256998 AL047055 AA489611 AA325311 AA374272 AA191652 AW068352 AA192283 BE265168 AW875505 AW401551 H64734 BE082132 N28721 BE270502 BE542010 AA095724 AA343838 AW862085 BE618237 N87961 AW402423 AW751453 AV648593 AA181533 W22134 A8184282 W27728 AA093676 BE620919 AA190693 BE618676 AW381000 Al675188 AA641654 AV648830 AV649121 W28254 AA133363 BE270651 BE314938 AA305540 W45064 BE613301 BE612706 AA774700 AW384452 Al129236 AA057759 Al904807 W26745 AW084436 AA092287 AA001711 AA148070 AA178879 BE614378 AA179254 AW673499 AW579754 W28174 AA083731 AA329644
15			AA491628 AA112012 AW673497 R57285 AW890681 BE614541 BE613168 BE184013 Al564430 T20087 AA357545 AA370341 AW020002 BE145977 AW386319 Al133645 AA928522 AA320082 Al564564 BE146370 BE535868 Al689539 AW364263 AW024767 BE181237 Al567727 AW386202 Al571579 AW007432 AW384430 AW751655 AW062554 AW190747 W25883 AW935205 BE439825 Al814738 Al609535 AA091211 AA847334 W25852 Al676217 Al917915 AA133283 AW385898 AW363381 AA676500 Al609403 Al922710 AW364148 Al819125 Al813788 BE439591 AW440632 Al669641 AW607753 Al961251 AW170466 Al813789 AW152388 AA733181 AW881158 Al829362 Al954017 AW089722
20			A1922140 BE613363 AW168114 AW471245 A1244988 BE612632 BE270925 W23836 AW753365 A1560061 AW607746 AA577685 AA856957 AL035899 N95408 BE546918 AW248541 A1889518 W24244 BE439756 AL493263 AA522799 T12329 AA211606 AW994980 BE140459 AA088784 AW948548 AW473607 AA191367 AW518727 AW882031 AW995169 AW995230 BE350742 AW265117 AW068516 A1590933 AW675814 W44616 AW675815 AW069302 AA088433 AA563768 AW805583 N30472 AW770831 AA055432 A1185055 AA121701 AW361826 N69778 AA192372 H05914 AT719090 R93535 AA129531 AW675529 AA595638 A1922588 A1422336 A1272104 AW151094 AW770433 A459624 A169786 AW805563 A16978 AND AW6756 AND AW6756 AW80756 AW
25			AA618622 AA258849 AW069346 W94137 AA120950 AW117291 W45032 AI660716 AI075409 W94042 W95255 AW368447 T92573 AI753582 W61154 AA351993 T92645 AW151115 AA022712 W61202 BE547553 R93536 AA988264 AA384897 AI094678 AA022677 AA092958 AA356987 BE301717 W44597 BE539272 AI298981 W28476 AA857019 AA176406 8A266 AA364932 AI376216 AI697013 AA176377 AA047415 AI368827 AI149318 T29905 AA223193 AW516507 AI952675 AA179002 AI523190 AI783877 AW473510 AW591026 AI689234 AI344434 AI299568 W45074 AW591339 BE042704 AI625502 AI523214 AI130808 AI719319 AA577323 AW473063 AW078789 AW068499 AA143226 AA527432 AW275892 AA100109 AA508682 AA566859 W95157 AI300997 AA534023 AI128750 AW386900 AA614319 AA962754
30			AA856565 A1000278 AA653314 AA779547 AA206056 AW750900 D52315 AW474584 N28645 AW276688 AA838489 Al282260 BE139186 Al619548 AW779619 Al476382 Al037996 AW263016 Al250703 N23056 Al285729 AW578650 Al289903 AA969812 AA486870 AA320509 Al560614 Al864164 AA353053 AW243842 Al358342 Al797438 Al752952 N94562 Al752940 Al700314 AA057760 Al535686 Al535709 Al535748 AA599435 AA676481 AA528726 AW272953 AA482791 Al077656 AW939918 AA577529 AA506613 H65225 Al700337 AA381152 Al719010
35			BE073480 BE073615 AW673514 Al289146 AW938944 AW939006 AW938996 AA56484 AW602284 AW938978 AW938935 BE073538 BE162787 BE073576 BE073462 Al459815 AA953334 AA716636 Al950483 AW779608 AW938951 Al889673 BE073553 BE162774 BE185856 AW602273 BE073474 BE073483 BE073618 AW602281 BE073542 BE073475 AW939008 BE073556 AW840704 BE073489 BE073463 BE162767 BE073562 BE073470 BE073496 AW602274 AW939015 BE073512 AW602307 AW602314 AW602294 BE073568 BE073557 AA497029 BE073548 BE073511 BE073573 BE073566 AW939039 AW939026 BE073513 BE073466 BE073484 BE073493 AW603154 BE073466 BE073514 AR05020 AR7444 AA56504 AR0502 BE073519
40			BE073625 BE073464 AA826629 AI721117 AA350594 T18953 BE007495 AW386898 AW793511 BE049519 AA565171 AW615051 AI347369 F01518 AI453289 AW071021 AW903128 AI718581 AI865531 AI753430 AW195177 AI753768 AI073727 AA599422 AW581743 AA657963 F01838 AW873884 AW613604 AW118006 AI521539 AW189876 AA853434 AW078573 AW027822 AI949090 AI753184 AI350844 BE162788 AW579284 F21714 BE156356 AI933613 AA953358 BE162875 BE070057 AI890891 AI688843 AW410101 AW276840 AA587013 BE070108 F29773 AI282914 BE070061 AW386904 AW386916 AW194231 AI446479 AA953634 BE161417 AW581762 AW581777 AW581774 BE161422 AW050520 BE543373 AI355481 AW386910 BE547892 BE544480 AW079762 AA528582 AA665889 AA148030 AI253726 AI253505 AW366859
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50	131524	36097_1	R96766 Al907845 BE142664 AB040927 AW503387 AA044786 Al686957 AW157364 AW976667 AI752687 AA191323 AA568170 Al161414 R17699 Al140787 Al140789 Al140788 AA742642 AA044809 AA485805 AA847859 AA480178 Z45709 AW974554 BE043079 AA809758 AA648838 N46563 AA485676 AW304745 AV657192 Al553650 AW118847 Al871278 BE075093 Al243817 BE046860 Al560949 Al669278 AA860508 N39152 AW131465 AA767854 Al457964 AA906227 AA719622 Z41372 T93491 AA954262 BE537985 T96329
55	101461	14616_1	N98569 AA029143 W15554 NM_000300 M22430 T39452 T47319 AA371017 N57336 D58694 AA320723 AA296089 R50467 AI346657 BED71861 AA319881 AW843848 AA320114 AW820896 AW610375 AW3933463 AW179006 AW610166 M22431 T46945 R70570 H02725 AW055209 R63131 AW392662 R86611 R53758 AA885780 AI749547 AW391366 AI970800 AW610360 AW821761 AI830923 AW000798 AA320004 AI249110 AI720962 AA682561 AA586608 AA643641 H00742 AA582755 AA609109 AA370548 R53759 AI672244 AI832430 AA683552 AA838623 R64075 AA838945 AI445267 AA534281 AI149280 AI274363 AI698468 AI128751 AA707159 AI150776 AA131825
60			Al991026 Al248667 R77118 Al168206 R50468 AW051904 N80785 R77117 AA037587 R63969 N32242 AW470160 R80612 R22811 AA320457 Al734854 AA534110 AA770261 N69947 R63087 H02619 AW768748 AA507524 T53621 AA535170 Al201371 BE439722 T61271 AA554850 N69967 AI805226 AA320357 H00653 AA229266 AA029021 R25199 Al659785 Al784636 Al186301 Al916742 Al581906 Al924395 Al569357 Al582097 AA534166 AA5333431 AA758464 AA642546 AA593596 R70480 AA327298 R21194 R24705 N98328 AA131992 AW150658 AA533307 T29484 Al198476 Al581608
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15	•		AW675475 N33466 AA844406 Al338233 AA804279 Al140172 R08039 Al371372 AW779420 AA973959 Al350202 Al311858 AA844068 AW051465 R58917 AA262562 AA255513 AA179827 R43808 R43783 AA179748 AA160554 T85488 AA223203 N70089 R52121 R43981 Al350079 H24363 AA844320 H80996 H72681 AA281259 AA863323 AA746986 R42662 T85278 AA194361 Al770137 F10117 T89022 F02675 AA194370 AA287013 N69852 T40917 N48265 R36686 H06305 T29294 R46268 H69825 R45142 AA166673 AA910318 R06282 AA203747 AA194455 R08090 R09678 F01096 R00610 R06340 AV651346 AV651308 AA089496 R57012 BE276300 H22856 AW751492 AW751491 T66408
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65			BE019193 NM, 005412 U23143 BE019174 H96758 L11932 BE294128 H39749 AW411386 H96748 AA131652 BE398084 W19483 AA747352 AA377688 BE560803 AA410805 R34798 AA323741 AA316917 BE396043 AA131143 AW410724 AL135023 BE294085 BE018423 AA361530 BE250748 BE299994 AA313516 Al653986 AA321590 Al591086 BE299410 T77128 AI951452 Al538143 Al609108 AA369495 AW411251 AW386906 AW082929 R95063 BE297935 BE019318 AW411277 Al691054 Al700140 AW474021 Al887587 BE439939 AI950991 Al625764
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	132349	28499_1	AW975654 AA652500 AW973307 AJ990990 AA918966 AA918970 AA612829 AW582941 AW584007 NM_003122 Y00705 AA983320 AJ660251 AW471481 AA845156 AW005713 AA551894 AA844948 AA586834 AA835291 AA627501 AJ302919 AA845077 AA921372 AA569123 AA974970
5	101600	21981_2	M20530 Al362622 Al310323 Al459618 AA919095 AA844953 Al362470 AA835382 AA740207 BE561617 W37399 AA405963 AA436100 BE393718 AA317393 C19033 C17096 AA365285 AA308297 AW410777 L10138 AA523678 AW328205 AA865267 R65897 W45653 N36027 Al065053 W44608 W44342 W919043 AA36178 AA379874 H38877 AA374479 AA151469 AA305168 C17593 AA305120 D55441 C17625 AA314253 AA211199 AA314589 AA310648 AW328687 AA021359 AA182541 BE565927 BE268825 AA706725 AA185325 AA187406 Al309641 AW006665 Al041213 AW327826 BE073571 Al610917 AW406432 AW600318 Al668800
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25			BE268132 BE391173 BE268771 BE513509 BE263242 BE276783 D19628 AW965243 W90258 AW962703 BE407481 AW406844 BE253043 BE073635 W15620 W47076 W78044 BE259558 BE565194 R00158 A434559 BE567338 N31298 BE409711 AI752598 AA100988 AA582094 AA902336 BE568220 AA167302 AA018050 AA915972 AA609370 A1184035 AA405697 AI208311 AI536822 AI583135 AI025630 R55812 R5507 AA987581 AA405807 AA082502 T40385 N88454 AA643754 AA093458 R85775 C14737 N24736 W31655 W47630 AA927763 BE268452 W73715 AI984453 H51401 BE537021
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65			BE465326 BE503131 Al670792 Al685623 R41236 BE348349 Al439167 AW469600 AW172580 R39862 AW130015 Al040222 AA609793 Al753801 R88603 Al288562 BE043005 Al693132 AA160883 H09133 Al684835 N57440 W51882 AA482283 AW505289 Al800765 H97982 Al807890 AA156810 AA156842 AA085507 AA085490 BE348821 Al932782 Al093768 AA280417 AA214471 T23573 Al857238 AW612154 Al351946 N35165 T32021 T31992 T32027 T32022 AA894475 AA877033 AA937480 AW952613 Al285827 AA810590 AW513250 W39699
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5			AW901191 Al268182 H19403 AA121223 D52401 D52407 D54786 D55281 W29029 Y00691 AA663488 M85440 AW246120 AW166102 AA325339 AA365967 AA422074 Al089418 Al858993 Al934500 AA808119 AW498597 W21815 AW157538 Al498264 AW157062 AA312480 AA450189 Z20539 W23223 AA707786 Al290619 AL133823 Al819350 N45630 AA323692 H44142 AA324278 Al829283 AW615583 AA131334 AA192364 AA129662 AA318854 AA319480 A1767204 AA233642 AW516760 H25864 Al961923 AA928093 Al923661 N50040 AW090665 BE300899 AW471178 AA722936 AW970337 AA533963 AA969003 AA703706 AA6687474 Al885747 AW003621 AA988680 AA708466 AA709132 AW872840 AA664299 AA505601 AW249567 BE504375 Al955868 AA811574 AA702297 N26094 Al361920 AA723293 AA019718 H46682 AA971391 AA709459 Al002552 AA302335 H20496 AA987656 AA668640 N92267 AA010292 Al052757 Al538598 AW150310 AA064742
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10			Al739170 H67915 Al304569 AA926749 AA031857 Al266691 AA938649 AA618497 AA723513 AA596001 AA470895 R73113 AA135724 AA703615 Al282652 AA865014 AA554895 Al076871 AA029470 Al791608 AA953956 AA523248 AW797855 Al567853 AA723757 AA228918 AA229626 Al688392 Al868189 Al202062 W42762 AA876330 R26067 AA548143 AA044388 AW272276 N56172 R54763 R70265 R23061 R23472 AV646146 R62911 T56208 T52223 AV661866 AA247611
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5			AW374302 H23317 AW176438 R07232 AW849553 AA352210 AW849234 H4665 W94208 AV653813 AA332023 AW250706 AA102094 AA321563 H29164 AW277021 R55980 AA355314 AA360928 N76776 W48614 AA026281 N23771 AW377275 D53282 N28452 H75676 D54754 D55368 D55402 BE181975 F07893 AA355322 BE566904 AA148259 T19343 AA193069 AA083979 AW404668 AA131541 R02406 R08392 AI669055 BE174166 T99624 AA985358 AA985342 AA131697 AW246850 W63788 AW601918 AW074311 AI951130 AI991110 BE070315 AA151763 AI032815 AA613379 AW182204 AW265712 AA747801 AI190719 AA430164 AI94088 AW131526 AW072023 AI004081 AI144397
10			Al301484 Al590672 AA970146 H99689 Al928778 AA928418 Al815800 Al816268 AW574757 AA102095 Al890190 AW340641 H62539 Al417393 Al608972 AL041908 Al872231 Al984699 AA317075 T99625 F24396 Al161392 AW162148 AW518979 R08341 H48776 F36841 W92663 AW161779 R83726 AW885483 AW440322 AA541604 Al908605 R59607 AW270365 R07180 AW273047 AW162487 Al052719 AW007925 AA366437 AW515604 AA206134 AA322919 AA836608 AW571844 AW023321 AW732262 AA220926 AA405177 AA564787 AA364610 BE220511 T71979 AW157224 H12389 Al990979 T72210 AA857354 AA292662 AA625743 T87002 AW058112 Al312077 AW969444 Al218167
15			AA534735 AJ908780 AA401454 AA947906 AW406988 AA864810 AA483282 AW170751 AW080972 AA830202 AJ151023 AA181703 AJ914655 AA889601 AJ905706 AJ201824 AJ141932 AJ680401 AJ283019 AJ365250 AI276334 AW410862 AJ056909 AW102782 AA807321 AJ185943 AJ920812 AJ140684 AJ378069 AW000855 N57455 N27435 AJ741814 AA192920 AA024282 H43629 AJ183784 AA563831 AW377321 AJ084030 AA074123 AJ318047 AA053173 AA872727 AA133110 AJ929302 AA132757 AA232267 AA129857 AJ033785 AJ346414 AA148621 AJ925122 AA932952 R51330 H08557 AA083810 AA129337 R55981 AW002267 H99629 AA625742 N53963 AA772739 T29282 AJ859385 AA223244
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25			Al698328 AW473013 Al860131 Al679762 T92179 H27238 BE265616 D52299 BE615720 BE614759 BE545737 AA074684 AA089952 D53006 AA232397 AA148760 AA182533 AA232066 AA243199 AA017339 AA018779 N94178 AA071301 T72989 R11772 BE387096 AA773305 BE379275 AW512721 BE256922 AW849540 AA861806 BE250923 AA149746 Al672946 BE565676 Al249672 AW377375 AA380648 Al475401 R11842 AW160619 R02307 AA017065 T85654 D56357 AW367156 R36960 D54522 AA211025 N59047 AA095616 BE268345 Al950148 BE148896
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5			R24153 AA916818 AF115106 X76648 H54165 W52609 W79548 AA279279 AA321573 BE379914 AA037204 AA011591 AA319951 AA312373 AW404037 AA308644 AA343991 T12247 AI188699 T28674 AI417967 AA313657 AW950049 AA485178 R67530 D21238 AF162769 AA131943 AW009604 H53565 W47497 AI884791 AI569215 AA845422 AA133992 AI743284 AA130209 AA779484 AW803387 AW803251 AA576037 N40812 AI580888 AI242038 AA830604 AI000536 AI921820 AI688612 BE439978 R24017 W31100 R69362 R20935 R81374 T79523 T79522 T87471 AW571801 AI244756 AW978497 AA954264 H90449 H54008 AA291163 W44537 AA033593 N51182 R93917 AI127891 T79435 T79434 AW803388 AA933060 AI141910 AI025368 AI095562 AA682885 AI151130 AI537290 AA779362 AI339261 AI242537 AA703360 AA131944 AW390025 AW389998 R67531 AA996112 AA931084 AA676656 AI373155 N94318 AW803369 AI041788 T79958 AA594671
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15			Alga4804 AW581359 Al312592 AW839221 AA328705 AW176725 AA046543 AW840706 AA070776 Al560816 AW889297 AA693461 Al016493 AA373039 BE162918 Al564313 AW004826 Al569405 Al419439 AA665773 Al828468 Al956098 BE504589 Al767273 Al684182 AA293733 Al473804 AA483299 AA340821 AA873749 Al669519 AA045473 C06097 AA505504 AW104511 Al281389 AA678558 BE551784 AW183737 AA725001 Al222869 Al751121 Al498613 AA598955 AW511819 AA746094 Al783501 AA857548 AA460980 Al183689 AA223510 AA993812 AW608353 Al751857 Al272689 Al090084 Al291427 Al185072 Al341799 Al150420 Al624079 AW075585 AA223394 AW511803 AA665468 AA903413 Al752782 AA679727 Al752273 Al061274 Al049690 BE464029 Al369788 AW628681 AW889130 W52099 AA968750 AA121393
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25	103269 102632	24390_3 32834_1	AF230662 S79332 AB012575 X79200 N24445 N72228 H99655 AW291679 AA730715 AW975672 Al699644 U66618 NM_003077 BE244439 BE244967 BE242731 AW403224 BE273228 AL079523 AA580844 Al910318 AA121441 AW853822 AA355815 BE257890 BE254018 R54755 H72035 AA378378 AW846512 AW375588 AA402352 AW404560 A133585 AF113019 Al174740 BE383453 AW375602 W69245 AA323991 AW369719 Al392720 BE076070 BE076246 AA252195 H54149 AW351946 AW371236 BE093129 AW317069 AI816724 BE463581 Al076940 Al123801 BE328399 AI870273 AA603776 Al039302 L31997 W69140 Al040190 BE301142 Al392962 AW189391 AW473894 AW082757 AA605026 AW168194 BE466942 Al222903 AA122012 AW275193 AA654229 AA847182 Al688771 F27116 Al871518 BE221378 AW439462 AW189556 AW193491 AI471650 Al799779 AW242172 Al580420 BE502872 AW452211 AA639503 Al368655 AW192152
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35		0.002.	BE293575 H95308 Z82206 BE388670 BE272057 AA216717 BE394775 T70795 AA304678 H78320 AA371448 Al684596 N89414 Al806645 AW883131 Al983340 AW858827 BE154049 AA094511 AA304354 Al806153 Al708422 AW604518 Al765617 Al735380 AW075958 Al963624 Al740521 AA096216 Al459163 AW339112 R00712 Al589173 AA602549 Al87959 Al762362 Al937378 AA341466 AW517324 AW157415 AW675434 AA613727 AA576788 Al186410 AA837040 AA165458 Al460082 AA404279 AW004639 Al240300 AW183961 AA243348 AA906904 AA243703 Al275411 Al283440 AA406265 AW661954 AA947892 Al015132 AA977942 AA557310 Al206823 Al399554 Al241703
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5			BE168373 AW243296 AW082309 BE550935 AA128024 AA659473 AW510824 AA931858 AA128067 AA126620 AI681167 AI829020 AW341066 BE222447 AA126493 AI814350 AW629523 AI340243 AW514576 AI765186 AI332628 AI275546 AI805954 BE467620 BE502644 AA573225 AW271279 AA825364 AA478779 AI882592 AA573330 AI7031177 AI738630 AI362767 AI272834 AW614575 AW440959 AI988599 AI769123 AI027197 AW051644 BE465249 AI28527 AI890976 AI299686 AA968889 AI266976 AA935053 AA635761 AA824445 AI400172 AW770719 AA965094 AA975535 AW440812 AA236455 AI653261 AI636923 AA807615 AA971410 AI498399 AI373497 AA973830 AW235436 AI766962 AI253126 AI399784 H30380 AW469278 AW193471 AI493399 AI093994 AI926918 AI468555 AI300692 R60701 AI027391 AI656958 AW263066
			T58718 Al521175 Al094088 T32368 AA470382 AW802933 AW802931 Al392989 AA777751 Al926386 AW779202 Al612991 Z41574 Al868604 Al825569 AW590300 Al825092 AA845416
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25			H67966 N72440 H79590 H81459 H60508 R39623 H60900 H40547 AA377244 AA318430 H71201 R64651 R65629 H72546 AW798947 N76974 H03029 N77701 AW151751 H60925 AA455839 H72947 N58334 N55487 AI299891 AA581634 AV651323 AV651728 AV650086 AV651295 AV648042 AW020600 AI537887 AA429713 AW080244 N73463 AA471335 AW150316 AA360851 W01407 BE074301 W21371 T87221 AA648042 AW06091 D16906 AW862400 AV661466 AI357816 AA442743 AI189966 AW887793 BE005206 AI926016 AA317024 AA976151 AA247314 AI490691 D16906 AW862470 D57965 N74437 N74336 H60409 N66059 H91165 R79462 F09991 R26175 H77853 N32590 D56667 AA461122 D56666 D56903 AW021856 AA374084 R69734 H66894 T81638 T63958 W23935 R67668 AW021682 H81249 H61959 H89852 R79306 W25710
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5			T34515 AA173847 BE560244 AL079696 T34244 BE389128 T34110 AW249130 BE266743 T04892 BE264706 BE264738 AW403234 AW401535 AA456984 BE207185 AW247060 AW998463 AA121373 AW994780 AA126661 AW407083 AW379415 AW578239 BE018419 AA366688 AA608513 BE378292 BE378337 AW103935 BE397789 AA302580 Al690498 Al697283 AA403211 N85842 AA463406 AW006584 AI589068 AA780276 AI871938 AA626835 AW439624 AW272829 AW512621 AI183842 AI591229 AW337254 AW410814 AA608753 AW009146 AA303726 AA456910 Al627190 AA877724 AW337702 BE302029 Al265957 AA775202 AW248698 AW245062 AW250412 AA173796 AI554428 Al244134 AA976264 AW086156 AA587634 AA425288 AA121363 AI377104 T58642 AA812669 AA812844 AI142489 AW074600 AI149028 AI588887 AW249803 AA425377 AA548184 AI919273 AI363195 AI344708 W76340 AI818834 AA622965 AI568201 AA101545 N52934 AI049533
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5			AA804318 AW518376 AW385913 AW083674 AI434034 AA938397 AI568283 AI249444 AA429884 AI719349 AW242259 AA429853 AW083900 AI738855 AI918581 H52597 R96654 AA341390 AA293559 AI680255 AI366472 AI472682 AA676709 AI718177 AA164192 AI339296 AW797221 AW511915 AI589416 AA953338 AI500000 AA876313 AI620034 AI865454 AA659294 AW474126 AW518454 AI719715 AW591921 AW379873 AI224057 AW302596 AI540725 AI499983 AI865593 AA728993 AI982586 AI865084 AA635760 AA946758 AW103642 AA742526 AA343052 AW474659 AI468868 H82407 AA485278 AW518420 AI468940 AW801881 AI497706 AW798431 AW802047 AI865054 AW806331 AI865359
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10			ALTOSSA ANDOUSS WOOT OF WOOT OF WILST AND SEASON AND SE
15			AA853566 AI590633 W58088 AW014564 AI880250 AA344158 AI131070 W81074 AI049815 AI073348 W58435 AA227175 AA496550 AW168598 AA872429 AI190794 AA478028 AI193735 AA055562 AI537541 AI018639 AW136662 AA853014 AI565886 AI915736 N36240 BE350339 AI750746 AI090756 AI75314 AA853251 AI753168 AI150013 AI249338 AW779255 AW027669 AA852109 AA291980 H72094 AI473398 AA455219 AW196020 AA293319 AA411063 AI682585 AI916014 AA732770 AW083712 AA372104 AI952485 AA250766 T47329 AA454876 AA055831 AA853849 AW241683 AW516455 AI553637 AA151152 AA917756 AW166676 R38279 AW068143 AI186543 AI568846
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25	102968	24693_1	AW582668 AA045209 AU076611 BE243124 AA343955 AA102186 AA307928 AA227387 AA314740 N86980 X16396 NM_006636 BE544371 AA361582 AA353457 AA172125 AA362045 BE545634 AW935573 AW368435 BE569039 N83453 AA354875 AW504993 W31001 AA281820 AW365599 AW365610 AW365606 BE542972 AI871407 AA480994 AA406118 AW387154 AI638274 C16082 AA210794 AW073141 AW365614 AI393287 AA282659 AI423141 AW365598 AW241512 BE466661 AI037877 AW591887 AW188655 AI683945 AA480995 AW365577 BE090663 AA171977 AI762124 AW875574 AI423138 AW951169 AW134804 AI263972 N98720 AA137110 AI439485 AW608789 AA551270 AA362384 AI739051 AA362456
30			AND
35	102976	14633_1	AU077174 BE616323 W25010 AL110099 NM_004390 X16832 AA305392 AW391441 R35036 AA309061 AA360805 AW815440 AW815685 BE171565 F06637 R18814 R64213 R68725 X07549 W78193 AA336414 Y18461 AW581172 AA054490 W95582 AA487325 AA487346 AA279600 H96491 AW969271 AA345648 W68555 AA992941 BE563114 AW976109 A1096690 A1122617 AI749481 AW026323 H69902 A1886820 T62554 AA283118 A1198774 N95617 W00965 AA593005 BE222876 T63395 A1004598 T91528 A1582812 A1126013 A1937274 A1031991 A1312045 AA844465 AA743166 A1872897 AA677296 W94703 A1380769 AA843780 AA908172 AA954631 AA864193 AA810948
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45			AA838328 AA658289 AI362589 AA872881 AA470832 AA659748 AA058337 AW512820 AA644662 AA502165 AI720562 AA865348 AA621446 AI267923 T59931 AA864686 AA582752 AA782286 AA768026 AW021729 R49423 AA622952 AA627103 AA913103 AI565188 AI869711 AI335624 AI363105 F02909 AI123813 AI919432 AI560364 AW591804 BE243287 A5540 AW518413 AI142065 T29991 AW963949 C02398 W70329 AI951124 AA747317 AI205593 AA947141 R64116 H73830 AA587498 AA532564 AW815511 AW815773 AW815836 AW815686 AW815435 AW815441 AW815775 AW815826 BE254396
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55			AA772163 Al076246 AA368689 Al247220 AA643529 Al240585 Al870488 Al075769 AA125857 W61147 R06238 H30656 AA159412 H30218 R01022 AA074396 AA355706 BE271450 AW157537 Al830027 Al816134 Al825881 Al688335 Al214071 AW176662 AW732611 BE562933 Al869702 AW997321 BE149700 AW821680 AA627389 AW856162 AW840172 AW856160 AW856164 AW856053 AW840205 AW582458 AW856165 AA340614 AW452775 AW382070 Al032419 BE272692 AW816677 AW816678 BE271792 AA576444 H67217 BE177808 AW816622
60			Al400391 BE149697 AA594450 Alw935536 Aw376579 Aw364201 Aw362605 Aw364193 Aw846801 Aw376626 Aw947474 Aw376794 AW250260 Aw376808 Aw376722 Aw376787 AA359700 Aw848535 Aw37671 BE143111 Aw848737 Aw848747 Aw848873 Aw376652 Aw848530 Aw849127 Aw848641 Aw848531 Aw848530 Aw848530 Aw848194 Aw848682 Aw848669 Aw848541 Aw848601 Aw848394 Aw376710 Aw848532 Aw849125 Aw376723 Aw376835 Aw848605 Aw376535 Aw848664 Aw848684 Aw848944 Aw348685 Aw849941 Aw376648 Aw8489198 Aw376608 Aw848200 Aw376589 Aw848554 Aw848879 Aw849095 Aw848571 Aw376602 Aw578344 Aw848941 Aw376648 Aw848198 Aw376608 Aw848200 Aw376589 Aw848554 Aw848879
65			AW848323 AW578331 AW848330 AW848319 AW578377 AW849088 AW376793 AW376669 AW578352 AW849126 AW376727 AW848594 AW848251 AW848241 AW578333 AW848434 AW578332 AW848913 AW848545 AW376574 AW848545 AW3765791 AW578335 AW848666 AW376676 AW376666 AW3766604 AW848529 AW578330 AW578338 AW848524 AW848517 AW376709 AW376634 AW376525 AW376763 AW578350 AW848524 AW376760 AW376670 AW3766797 AW376755 AW376773 AW578350 AW848258 AW376760 AW376603 AW848446 AW849139 AW848611 AW376761 AW376611 AW848887 AW376628 AW376716 AW848647 AW376703 AW376714 AW376594 AW376596 AW848457 AW376672 AW848190
70			AW346606 AW376586 AW848462 AW376742 AW376677 AW376598 AW376746 AW848610 AW848447 AW848683 AW376556 AW856261 AW376638 AW376538 AW367077 AA068995 AW068772 AL120506 AL120386 Al244494 BE545234 AA159304 AW376599 BE295933 AF187522 AA152400 BE543706 AW576337 AF188896 AA811996 AF189213 AF188897 AW856185 AF190059 BE070163 AA251742 AI750309 BE150151 BE150252 AW848663 BE150159 AW856128 BE150201 AW578313 BE150250 AW381440 AW856074 AW381435 BE150190 AW856179 AW856121 AW856174 AW401844 BE003225 AW856177 AA526169 AA640192 BE314916 AA779712 AI906402 BE537879 AI834242 AI909763 AW845219
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80			R34799 F11143 A198270 AA335689 AW439092 AA577609 AW894217 Al302960 AW406637 C17923 H63348 Al634226 BE177956 Al498384 AW609479 AW950912 BE085889 AW391004 R14402 Al669187 AI758210 AW150328 AW402978 AW474568 AW579293 AW363558 AW369322 AA633069 AW364214 AA557144 AA352699 AW369361 Al652770 AA037067 AW369367 AW369378 AW369383 AW369320 AW369340 AW369334 AW747900 Al452805 AA025994 AW969302 Al471469 AW838332 AA700483 AW575707 AW363552 AW754279 Al538596 N41444 AW369369 AA533573 Al697373 N91447 T63645 AA343413 AW369372 AA329807 AA847288 AW369363 AA009432
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102903	5			BE177210 H43742 AA130321 H05246 AW517890 AW511678 AW198136 AA411576 AW934870 AW131725 N99045 AW190050 W48791 R96149 AW298454 AW051778 AI423040 AA147092 AW438903 AW519147 AI936035 AW747910 AI588900 AI572603 AA854132 AW272152 AI884403 AI689595 AA994684 AA576990 AW837159 AA411440 Z20745 AW369342 AI554272 AW051768 AA865624 AW190197 AW129438 AI590389 AI954048 AI553828 AI000547 AI858437 AA554141 AI017045 U82777 AA431097 AA622202 AA101026 AA890524 AI141907 AA770195 BE000705 AI338886 D20039 BE001103 BE000729 AI093763 AI128438 AI038158 AI014806 AI206804 AI128741 AA731344 AA781650 AA847245 AW977294 AI149018 AA725251 AU073592 AA601940 AA687609 AW001981 AA857855 AA027254 AI696346 AW068510 AA025935 AI288581 AA312064 AI366687 AI123208 AI361102 AA009433 AI375745 AI190304 AI810395 AI061262 AA7222843 AA027255 AA693398 AA983511 AI523184 R55808 AI766707 AA723485 AA158005 AA731346 AI860056 AA470452 AI344375 AI052053 AA706704
## ASSEDIES PAIL SERVICE SERVI	10	102002	17554 4	AW082751 Al354629 AA843429 Al701333 AA161092 AA226840 AA993389 AW176551 AA243693 AA937997 AA470742 R96150 AA873311 AA282110 Al245104 AA318159 Al648622 T63845 AA972595 AW573031 H63268 C75028 AA523040 F02163 Al221319 F04541 AA708486 F08814 AA550863 AA524127 T28847 Al868107 AA635688 AA282111 Al147151
112088	15			AA315209 NM_001237 X51688 AA158802 X68303 AA360411 AA001329 AW608728 Al061440 AW875571 Al654232 AW371180 AA608568 AW371208 AA213393 AA306347 Al872410 AA936671 Al763348 Al948484 N41638 AA482297 Al827243 AW276578 Al199011 Al350965 AA158803 Al040688 AA693660 T28292 AW950496 AA001916 AA213394 AA557629 Al872826 BE564910 AA580754 AA459213 AA213538
### APASSESSE AADSSESS WISSESS WIRGESS ANGESSES		112068	611385_1	Al264847 R43910 AW614197 Al863821 AW467620 Al695292 Al672346 Al302090 NB1071 Al611641 AW166600 Al168293 Al313201 R43835
113936		104204	11607_1	AA285262 AA055428 W52943 W78060 AI669713 AI804895 AI056890 AI202008 BE504324 AI638488 AI991279 AW301184 AI990138 AI765837 AI523554 AI735158 AI637794 AA922055 AW069634 AW875295 AW002630 AI089420 AA535017 AI652587 AI657071 AI637803 AA677262 AA865617 AI69986 AA223477 AA554162 AW606040 AI078073 AA573096 AI057436 AI307113 AI983310 AA723619 AI659825 AW275484 AA552067 AW134930 AI038417 AI247714 AI678270 BE139653 AI814032 AI424176 AW874195 AA234118 AA843211 AW136280 AI468611
104254	25			R44714 AI952898 AI623118 AI271632 T10160 Z40968 D86962 Z43779 AA298247 AF073378 AF000017 BE386788 BE146000 AL046008 AW951300 BE328763 AA565135 T17443 AW197239 Z39844
BE264316 AA358271 BE253249 BE621112 BE256248 BE253588 AF008557 M49006 AV4005919 AA352701 T25793 H51697 AB29974 W89006	30			W94015 AW518912 AW385139 N92349 F28925 C15300 Al239534 Al358889 Al625560 Al936054 Al239416 D60096 AA808160 AA889642 Al360831 AW197699 AA136336 AA807872 H92911 Al123784 W72779 Al146976 Al023919 Al183855 AA298388 R85083 AW085113 R35681
## 19451   16763_1	35	104254	28286_1	BE264315 AA356271 BE253249 BE621112 BE256245 BE253568 AF008552 N49806 AW4058 AA352701 T25793 H51697 Al829974 W89058 AA297276 AW170452 AW249668 BE392539 AA070238 H81023 AW339856 N49700 AW406366 AA071486 AA720659 AA847804 AW411426
4.0	55	134351	16763 1	BE045492 AW294622 AA883408 H82885 R97912 AA810605 W88963 H81024 AW103189 AI453120 AA738386 T25124 AI582910 AI352050 AI802294 AI348029 BE561223 H57656 AW407129 R97911 AW951023
A9311301 13142219 A3863447 A4774153 1026995 AA426099 W74093 AA001637 N69444 AA902567 R01486 AA639804 AA005772 AW0050972 H7327 AW73992 AA75927 AA00680313 AA43694 R22446 H72073 C00094 D29190 AA918874 AA577952 AW099231 N53786 T99746 AA586967 H00564 AA001256 H01095 AW76474 R70643 T63362 H03335 AA487014 A1346925 AW7272865 AA827933 AW0372012 AA917823 AA904510 AW151183 AW004410 T878777 A774429 T170503 T53333 W80029 W52504 AW074106 AW571586 AA335556 AJ355538 AJ922244 AW276403 W52888 AA588801 AJ433345 R82074 AJ343474 AJ893962 H95157 AA002260 A890064 EE049826 AZ006356 AZ01674 AW610406 AJ952099 AJ014897 R00530 EC049407 AJ820005 A422564 AW0711267 AW054439 AJ880402 T58571 AA568850 R015468 AZ022671 AJ864071 AJ955099 AJ43988 T11663 W54682 AJ678237 H3905 AJ49398 T17663 W54682 AJ678237 H3905 AJ49398 AJ8920 H395106 AA99862 AA502822 AA502822 AA502822 AA502827 AA50291 AJ49398 AJ49389 T17663 W64682 AJ67823 AJ8930 AJ8939			_	AA219419 AW374657 BE081779 AW352196 AW602851 AA368110 BE078507 AA299561 AA377906 T58819 BE273643 BE541572 AA367994 AA100094 N93982 R22500 AA375599 AW998547 AW887074 AW631259 AW085777 Al660836 H13083 D58798 AA010621 T39490 AA001108 AA224261 AA366381 AI696816 W79383 N4011 TA AI219172 AW630029 AW079051 AI829106 AW439517 AI814283 AA579623 AW084866 AW170078 T92786 AI860472 N54556 AW009667 AI333283 AI348031 AA707206 AI831036 AA928681 AW337157 BE160976 AI422988 AA777013 AI691025 AA032042 AI831457 AI921282 T89395 T62508 AI566209 AW516825 AI758659 AI271852 AI677918 H01094 AI829280 AI224622 D59025 AA723113 AA601514 AW192078 AI224154 AI015641 AW182754 R68745 AA031960 AA583770 AA921870 AA632080
	43			AA911301 Al342919 AA863447 Al474153 N26996 AA426099 W74093 AA001637 N69444 AA902587 R01486 AA639804 AA057472 AW050972 H13287 AW273894 Al872681 T47346 Al865585 T29391 AA745902 AA069313 AA443694 R22448 H70273 C00094 D29190 AA918847 AA577952 AW999231 N53786 T97945 AA586967 H00654 AA001255 H01095 AW376447 R70643 T53352 H03335 AA487014 Al346925
N23362 AA682872 H03110 A168530 N32346 T94740 AA236561 AA236235 N23955 R27720 N31614 A1814425 A1804857 AW590744 A1080155 N30796 A1341754 A1367163 A1272814 A1332944 AA193683 A1183991 239979 F04878 A1868457 N26707 BE535358 R23737 AW449959				W52504 AW074106 AW571586 AA335556 Al355538 Al922244 AW276403 W52688 AA588801 Al493346 R82074 Al343474 Al693962 H95157 AA002260 Al590864 BE049626 Al206365 Al241074 Al610408 Al952089 Al014897 R00830 BE049407 Al820005 Al422564 AW511287 AW591439 Al864028 T58751 AA568360 R81546 Al282671 Al684071 Al950509 Al439380 T71663 W94662 Al678329 T49501 Al079708 AA995106 AA918622 AA502982 AA502632 AA548291 Al354395 AW000944 AW085741 D45578 Al826443 Al810939 Al301212 Al243066
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N40523 AW474740 AA022464 AA993528 N46572 AA129732 AJ342643 AJ144408 AA011376 T95293 AA057170 AW613713 AJ280406 N95765     AA031474 AJ128171 AJ097021 AJ684137 AJ889755 N29991 H89564 W92688 AJ953558 H98678 AJ168616 W47201 AJ6890716 AA834490     AJ160430 AW207161 AA677837 AW080654 AW104712 AJ688138 AW474712 AJ559164 H89565 AJ85000 R76553 AA662930 AJ340202     AA608802 A757215 AA595069 AJ199506 AJ765271 AA028027 AA703651 AA918632 AJ128696 AA903074 AJ027793 AW204001 AJ932695     AA846139 AJ859558 AJ537176 AJ040586 AJ270245 T40492 AA987460 AJ160028 W23529 AA029035 AJ827556 AA904875 AA706779 AA807465     AJ3731	60	134369	5386_1	AA677116 AA368429 C03600 H2712\(\bar{0}\) NBB341 Al126019 AA373718 AW964293 AA345812 AW967361 BE047207 Al571069 Al335849 Al537518 AW168050 W47316 AL355724 R13547 Z43925 Al769318 R19976 W35345 W24878 AW194129 T95373 W07142 H28325 AA634915 T86778 T39243 T10738 Al370696 T36271 AW954313 AA296523 AA029247 Al369552 T40309 AA022997 AA011364 T41144 T41173 C18560 N59612 AA328867 AA040690 AA088617 AW167394 AA028018 Al569560 AW195344 Al089584 AA151507 AA133346 Al692832 R20636 AA031616
A846139 Al859558 Al537176 Al040586 Al270245 T40492 AA987460 Al160028 W23529 AA029035 Al827556 AA904875 AA705779 AA807465 Al689182 AW050514 AW150550 AW150472 Al648649 AA781059 Z41664 AA897320 AW050517 R45078 Al933450 Al740792 Al949422 AL079298 Al423046 N31952 AW195192 AV660395 BE543143 AA658226 R89511 R88931 Al740792 Al949422 AL079298 Al423046 N31952 AW195192 AV660395 BE543143 AA658226 R89511 R88931 Al740792 Al949422 AL079298 Al423046 N31952 AW195192 AV660395 BE543143 AA658226 R89511 R88931 Al740792 Al949422 AL079298 Al423046 N31952 AW195192 AV660395 BE543143 AA658226 R89511 R88931 Al740792 Al949422 AL079298 Al423046 N31952 AW195192 AV660395 BE543143 AA658226 R89511 R88931 Al740792 Al949422 AL079298 Al423046 N31952 AW195192 AV660395 BE543143 AA658226 R89511 R893953 H93849 R83420 R97372 H74255 H73034 R92870 H508 R491252 H62329 H50974 R73807 R26201 H50967 H81482 H78630 N94123 H69920 R89892 AA340135 R97379 W03876 H19610 T58178 N78051 W25742 H78463 R63791 H63594 R83842 AF097635 H73826 R94380 H47962 T53050 T54718 R64416 R74107 AF147332 T53241 H01120 R69900 R63176 R78653 H81168 R80097 R66776 H66298 H60727 R87144 R99227 T67143 R89448 H52368 H50753 R76757 W01513 T59136 R70829 N77678 R80500 H78988 Z84721 N77683 N49814 N49427 AL038057 T52784 AA176749 N74739 W03863 T56842 N73036 T48014 N58317 N23926 AA766008 AA054580 N27635 R24279 AA458708 T51891 R34338 N63760 T58643 T54776 T56870 T52464 R26296 R48636 R64108 T52509 Al266020 T56462 T52644 AW950562 T54777 T50483 T54790 T53061 N49288 T51607 R52129 T50474 H65397 T55568 H93336 H00452 R81632 N71325 T52168 N71376 A497856 R39511 R80225 T50668  80  T52574 AA069455 T52166 R28311 H68765 R89496 AA182860 T54943 T56607 T52370 R68553 H73383 H94478 N74134 T56851 Z20477	65			N40523 AW474740 AA022464 AA993528 N46572 AA129732 AI342643 AI144408 AA011376 T95293 AA057170 AW613713 AI280406 N95765 AA031474 AI128171 AI097021 AI684137 AI889755 N29991 H89564 W92688 AI953558 H98678 AI168616 W47201 AI690716 AA834490 AI160430 AW207161 AA677837 AW080654 AW104712 AI368138 AW474712 AI559164 H89565 AI185000 R76553 AA662930 AI340202
H74255 H73034 R92870 H50844 R91252 H82329 H50974 R73807 R26201 H50957 H81482 H78630 N94123 H69920 R89892 AA340135 R97379 W03876 H19610 T58178 N78051 W25742 H78463 R63791 H63594 R83842 AF097635 H73826 R94380 H47962 T53050 T54718 R64416 R74107 AF147332 T53241 H01120 R69900 R63176 R78653 H81168 R80097 R66776 H66298 H60772 R87144 R99227 T67143 R99448 H52368 H50753 R76757 W01513 T59136 R70829 N77678 R80500 H78988 Z84721 N77683 N49814 N49427 AL038057 T52784 AA176749 N74739 W03863 T56842 N73036 T48014 N58317 N23926 AA766008 AA054580 N27635 R24279 AA458708 T51591 R34838 N63760 T58643 T54776 T56870 T52464 R26296 R48863 R64108 T52509 A1266020 T56462 T25644 AV950562 T5477 T50483 T54790 T53061 N49288 T51607 R52129 T50474 H65397 T55568 H93336 H00452 R81632 N71325 T52168 N71376 A497856 R39511 R80225 T50668 T52574 AA069455 T52166 R28311 H68765 R89496 AA182860 T54943 T56607 T52370 R68553 H73383 H94478 N74134 T56851 Z20477	70			AA846139 Al859558 Al537176 Al040586 Al270245 T40492 AA987460 Al160028 W23529 AA029035 Al827556 AA904875 AA706779 AA807465 Al689182 AW050514 AW150550 AW150472 Al648649 AA781059 Z41664 AA897320 AW050517 R45078 Al933450 Al740792 Al949422 AL079298 Al423046 N31952 AW195192 AV660395 BE543143 AA658285 R89611 R88931
AA176749 N74739 W03863 T56842 N73036 T48014 N58317 N23926 AA766008 AA054580 N27635 R24279 AA458708 T51591 R34838 N63760 T58643 T54776 T56870 T52464 R26296 R48863 R64108 T52509 Al266020 T56462 T52644 AW950562 T54777 T50483 T54790 T53061 N49288 T51607 R52129 T50474 H65397 T55568 H93336 H00452 R81632 N71325 T52168 N71376 Al497856 R39511 R80225 T50668 T52574 AA069455 T52166 R28311 H68765 R89496 AA182860 T54943 T56607 T52370 R68553 H73383 H94478 N74134 T56851 Z20477	75	133731	33199_24	H74255 H73034 R92870 H50844 R91252 H82329 H50974 R73807 R26201 H50967 H81482 H78630 N94123 H69920 R89892 AA340135 R97379 W03876 H19610 T58178 N78051 W25742 H78463 R63791 H63594 R83842 AF097635 H73826 R94380 H47962 T53050 T54718 R64416 R74107 AF147332 T53241 H01120 R69900 R63176 R78653 H81168 R80097 R66776 H66298 H60727 R87144 R99227 T67143
	80			AA176749 N74739 W03863 T56842 N73036 T48014 N58317 N23926 AA766008 AA054580 N27635 R24279 AA458708 T51591 R34838 N63760 T58643 T54776 T56870 T52464 R26296 R48863 R64108 T52509 Al266020 T56462 T52644 AW950562 T54777 T50483 T54790 T53061 N49288 T51607 R52129 T50474 H65397 T55568 H93336 H00452 R81632 N71325 T52168 N71376 Al497856 R39511 R80225 T50668 T52574 AA069455 T52166 R28311 H68765 R89496 AA182860 T54943 T56607 T52370 R68553 H73383 H94478 N74134 T56851 Z20477

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10	103677 133797	41847_1 14537_1	AA454684 AA987257 Al351217 N92937 AA007661 AW389309 AA456333 T87606 D31579 D30835 T49124 BE042568 Al690934 T31165 AA843733 R86012 AA135350 AW389311 C03271 AA345798 R46788 NZ7045 AA995286 Al572405 Z83806 AJ132091 AJ132090 AL133921 BE389006 NM_005056 S66431 T07054 AW500214 AW604275 AA487706 AA211245 AA247515 AL133922 AA311252 AA487492 AA312860 AW268369 BE328608 AW105357 AW468600 BE535444 AW672876 U25911 AA877356 Al587632 Al609139 AW500785 AW997007
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	134444	33247_1	BE184455 BE396187 AL035660 NM_003064 X04470 AA132992 Al862145 Al564623 AA572950 AA993549 AA026099 AA460433 X04503 X04502 AA026192 AF114471 Al858387 Al885550 AW264225 Al638119 AA564454 Al222907 AA541595 AA587161 Al743345 BE044073 Al742512 AA551908 AW238407 BE392080 AA397776 AA863166 AA587140 Al042208 AA683520 R71834 AA026497 AW081599 AA932864 Al580185 AA316675 Al000873 AW513394 H65171 BE612494 BE612943 Al377093 AA938592 Al148713 AA594366 Al042358 Al282099
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75	133893	14927_1	A434699 X01060 NM_003234 BE256019 BE142860 BE142729 AW500605 AA347570 AL120908 AA082493 AW630459 AW501236 BE009458 AW503924 AA055688 AA216664 AW802703 AA877477 R82301 N85217 AA134422 H02417 AW389909 AW389907 AW389913 AW389877 AU46375 AA488721 R95492 A1189434 A1132910 BE092247 R11868 F05413 N85500 H60074 H02305 W95694 D59086 R82712 AW402489 A1630673 AA132188 AW629714 AA581142 R19476 AA033935 AW862307 F12939 BE536497 F12950 H13379 AW815251 AW861747 AA490726 AW861754 AW861737 AW366658 AW858160 AW858110 BE010851 A1630422 A1630188 AW852822 F12235 AW935238
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55	133908 133944	28620_2 5151_1	AU076820 M83216 AA971545 H51609 AA092764 AI926727 AI801609 AI888318 AW950682 R99241 N42334 T68396 R24753 AW083647 R01328 H50950 AW068579 AI205108 AL049969 AA249019 AW068578 AA056648 AA056482 R58113 AA056676 F13429 F11610 AW840189 AW948891
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20			Al765107 AK000557 AW250662 AW404558 AW631125 Al474992 H38495 BE259536 AA338808 AW961032 AW375527 AW375519 AA036978 T98407 AA058761 BE394031 AA366003 Al761506 AA587887 Al573291 Al744657 AA588536 AA738047 BE502073 AA700013 AW246799 T85297 AA759011 Al890594 Al148268 Al587531 T15960 AA830722 Al298775 Al200417 R00301 Al798469 Al825338 Al809308 Al436070 AW137267 Al344370 AA838214 Al085034 AA838498 Al537302 Al368683 Al040364 Al341279 Al365563 H78131 AA036979 AW082916 Al248063 AW337164 AW663937 AA683570 AW473394 Al500302 Al357177 Al927184 Al991231 AA365119 AA573353 AW630338 AW872754 Al766544 AW569264 T32118 AA719102 Al370730 Al469218 Al668957 Al263592 AW103958 AA434441 T99954
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30			AW305249 Al148765 Al244424 Al144495 AA279759 AA521064 AA907836 AA552043 Al000083 Al478931 Al128403 Al200568 H95707 F30237 Al302453 N94464 AA557186 F20977 Al491886 AW183938 AA569868 AA852663 AW275122 H91807 N41841 N30742 AW169997 Al038347 AA746065 AA972090 AA654533 AA654036 Al936891 Al807215 Al332537 AA906338 AA115690 Al251732 AW975227 AL046958 AA937615 AA730937 Al420622 AA978142 AW105551 AW182041 Al363204 AA651663 Al057284 N94563 Al827188 T90621 F18332 AA707791 AA635717 AW249748 N80826 T24920 F18565 Al005595 AW592487 Al674940 AA283903 AA084667 Al674878 T89384 AA552449 Al701784 F37912 AA215865 H91708 Al040250 N98724 Al264957 BE391925 N56092 AA369378 N56408 AA092677 F36194 AW023614 AA664664 AA665052
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45	128530	28930_1	AI932995 BE064464 AW371902 AW371841 AI885885 BE064457 AA524113 AA721037 AA504343 AA778099 AI800598 AI693112 AI864633 AI690228 AI400990 AW969089 AW371927 AW371912 AW383562 BE151089 AW383568 BE218503 AW383570 AW371899 BE151097 AW371900 AW293095 AW292008 AA434179 AA714780 R45868 W01182 AW957767 AW119223 AI207864 W01578 AA354403 AA805177 AI613299 AW269636 AA481528 AW079101 AF131777 R60489 T81289 AA481594 BE181020 AA465433 AW808125 T84992 AA749191
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5	105726	5801 <b>_</b> 1	NM_012068 AB021663 T71531 T67794 AA344893 H46645 AA191110 BE271163 AA513805 AA512936 T67718 T71368 D31104 AI870651 AW629156 BE207819 AA161164 AA292328 AA815137 AA994765 AA191099 AA994766 N59773 AI000315 H46624 W56638 T64935 W56601 R06908 N93059 T85073 AW390226 AA465295 H27386 R99387 H53559 AW971750 AA714781 T73635 AW381810 AW601287 AW601284 BE063948 AW601286 BE063943 BE064022 BE063949 BE063947 AV651606 BE063950 BE063954 AW601282 AA345127 AW601288
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14/292	5	107071	179431_1	
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105811 71767_1 1515178486 F28889 ABDIGAT W61382 AA027984 ADDISSOS T88374 AA237793 AXMS72381 H80399 AVT75147	15	113674	4406_1	Al985606 Al053516 Al312176 BE560905 NM_014214 BE407555 AW672705 BE538245 BE619341 AF014398 AF200432 AA054659 AA350997 R52482 BE257590 BE255436 BE540254 AA352378 AA368117 AA056721 N95677 Al742203 Al184977 Al703196 Al858501 AW190354 BE219864 AW468540 Al264208 AW088250 BE049259 BE222751 BE546756 W60101 AA297285 AW014869 AW675462 AW381525 AW513795 AW675783 Al624316 AA994724 AW468646
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60 105914 5785_1	55	113783	4882_2	AL359588 AK001821 AW160980 AW160713 R60610 R59877 H10278 AA344815 AA349679 AW937762 R51499 R20177 R20270 Al935430 N98574 AA557887 AA559968 H57311 AW957511 Al341683 H10222 R60556 N69972 R59878 Al015582 Al814829 AW515396 T33330 AA349678 Al635336 AW243924 Al371168 W19222 T17389 AW965984 R51500 Al358310 AW136265 F02645 R39158 Al269711 AW150587
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5			AL287542 AA632199 AA448204 AL198285 AL869256 AW574781 AL349346 AA713587 AL263944 AL864900 AL273251 AW515501 AL265988 AW780139 AA909952 AA765867 AA465092 AA744588 AA731158 AA814660 AA748723 AA768424 AA235743 AA765706 AA809967 AA632362 AA722465 AA721072 AI739477 AA806084 AL363026 AA252319 AA830765 AA768839 AA780188 AA737659 AA936522 AA251781 AA648959 AA402783 AA452833 AA262136 AA744737 AA458933 AA768945 AW016384 AL073667 AW140044 AL919463 W02122 AA227390 BE393943 AA282805 AL183859 AA830665 AL871369 AL887043 AA268895 AA594349 AA837324 AA214524 AA810962 AA287576 AA490613 AA825316 AA613373 AA761522 AA731427 AA767785 AA733191 AA948512 R32626 AA251671 AL860358 AL424667 AL493334 AA481488 AA252361
10			A731499 AA399475 AA737005 AW137326 AI824260 AA465210 AW247312 AW368898 AA302102 AA256564 AA912888 H24558 AA604467 AA252929 R23909 AI572501 AI858741 AI130884 AI538528 AI288244 AI991619 AI131338 AI379888 AI810139 H18623 AA523535 AW517344 AI279546 AI144242 AI379684 AA573326 AA4796596 N23081 AA397812 AA262774 AI186536 AI222012 AI192712 AI818719 AW071697 AI805329 AI871755 AI192609 BE220626 BE219396 AI633555 AW104172 AA577335 AI827811 AW007345 AA625830 AI588989 AW770682 AI148337 AI475084 AI123923 BE550812 AA662467 AI360115 AI805170 AI369687 AI193722 AI282176 AA972427 AA236915 AI36153 AI086426 AI222895 BE549696 AI123934 AI085689 AA235489 AI273579 AA564468 AI239466 AI298426 BE220634 AI417179 AI669072 AI187330 AA252247 AA434251 AI342328 AA477824 AI167911 AI417106 AA961041 AA642406 AA233129 AI082559 AA700492 H41441
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25	130094	26139_1	T28575 R10378 N92944 AF042036 T72498 AJ444654 NM_001471 AJ225028 AF099148 AJ012288 AL031983 AJ012185 Y11044 D80024 AL119755 X90543 AJ598214 X90542 AW380842 AW380854 AW380862 AW380861 AW380836 AW380767 AW380855 AW380765 AW380851 AW381524 AL042317 AA348199 H51356 H19658 Z44106 AW867915 BE181406 AW896205 AA181004 R71844 AW373750 AW373775 AW373673 AW373686 AW839098 AW373772 N56175 T81224 M78726 AJ741552 H43286 AJ371087 AW090617 AW264022 AW081493 AA348198 AJ668882 AJ435011 H19659 AJ078526 R76486 AJ808509 H41556 AA992062 AJ242458 R71794 AJ280436 AA742280 H50397 BE047768 AW517824 AJ866545 N70841 AW263371 AJ377521 AA908707
30	115084	10376_1	T81014 AW380864 AW391804 T99207 R73356 H25821 R76485 W01458 T99208 BE383668 AK001480 AA148764 BE612465 N29908 AI663707 AI803462 AI984336 AI244784 AI202881 AA101937 AA160974 AW672908 R70272 BE502676 AA076294 AI694452 BE466032 AI634396 AA127707 AA317157 AI431745 AI479348 AW298074 AI819738 AW779732 AI636201 AA127644 AI879529 AI929217 AW967042 AI979011 AW999434 AA160975 AI193979 AI261728 AI611182 AI220151 BE041823
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45	113803	37976_1	AV880709 AW29531 AJ20597 N05082 A9167 16 AA69210 AI57032 AND 1545
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55	113816 106639	437168_1 7493_1	H46088 AA757630 H45998 H47035 W45499 H18092 H46080 H46068 W40422 H18132 H46498 AV655272 Al382139 Al124646 AW298134 AA652260 T58540 Al337943 Al354941 AW511303 BE501483 Al371627 Al687503 Al693430 Al693871 BE348647 Al091164 AA947682 AA371477 Al014595 AK001478 AA313424 AA775305 AL119130 R13701 AA363659 AW959490 AA460066 T95465 Al161400 F07057 Z42134 AW298014 AA134238 H15216 R19551 AA356614 AW965786 Z43860 AA448444 AA133248 R09023 AA011707 W52631 BE539194 AA404459 BE540061 H77582 R65897 R82856 R77316 R07005 N76954 AA151044 AW237218 N45210
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5			AA228444 AA503911 AA565282 AA658335 AA574401 AA224941 AA652562 AA640888 AA652767 AA935442 AA641051 AA228914 AA934464 AA467738 AA468189 AA595856 AA635384 AA226579 AA657837 AA566022 AA659486 AA572841 AA658248 AA569582 AA639325 AA555126 AA229568 AA578333 AA533825 AA579236 AA508086 AI826974 AA658203 AA888477 AA228966 AA533715 AA886949 AA541337 AA888317 AA661941 AI969953 AA657879 AA550919 AA533343 AA657824 AA554865 AA687194 AA640720 AA569708 X75682 AA559285 AA632441 AA659520 AA533537 AI201973 AI400942 AI417483 AA528834 AI824643 AI805186 AA548801
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65	101013	31218_1	BE300094 BE384439 AW794648 NM_002305 M57678 Al929016 AU076727 ZB3844 ZB3844 Al906100 W44519 H98497 AA188069 AA572687 AA035793 W93978 BE409220 AA359751 AA502475 H28319 AA527889 AA432335 AA864762 AA340061 C05180 W68192 AA327811 AA345871 AI750205 N34093 N86639 AA085753 AA603451 AIA542262 N42135 C04367 N57266 AI038364 Al184846 Al928853 X15256 J04456 AA603552 AA317300 AA568615 AA813495 N40276 AA4002624 AW264898 H21418 AA643822 AA603569 AA507955 N44497 AI000869 AW079049 AA614629 AA303987 AA362817 H54502 N85495 W52256 F30575 AA568129 H26935 W93977 AA373651 AA872398
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80			AA953030 AA588476 AA131216 T79619 AI752885 AA614820 AA988962 AI143561 AA493182 AI302481 AA301613 R73520 AA069898 AA374944 AW364221 AA342013 AI244949 F36390 AW050980 N79486 AA101160 T68112 AI750204 AA328787 H02617 AA314734 AA527923 AA307835 AI885112 AI872905 AA534666 AA188363 AI192490 H45772 AI824700 AI184276 AW079473 N29847 AA720843 AA720914 AA573391 H54416 T59424 AI824457 AA304220 AA482553 W72882 AA627932 H27514 H28400 W68050 H20953 AA635786 H21376

5			AA514046 AI342823 F29905 H25999 AA757144 H21636 F22104 AA428650 F27143 F28346 AA535690 H45771 AA548851 AW170154 H45646 W92274 AI921614 AA176461 AW170153 AI927284 AI161206 AA594439 T28595 H41129 AI497579 AA978015 AA328875 AA373653 AA090973 AA328623 AA328759 AA366468 AA375406 H46976 R86050 H02722 AA328321 AA328205 R62358 AA373717 AA304138 AA304224 AA301603 H54867 AA374783 AA376232 AA373239 AA374917 AA375673 AA303857 AA376466 AA376461 AA302613 AA304082 AA301731 AA357988 AA303328 R25744 AA301587 N78746 H20508 AA659423 R47960 AA825456 AI001806 AI245114 AA7292223 AA860271 AI913845 H26296 AA733035 AA340965 AA304291 H27356 H20598 AA129613 R69996 AA157689 H20992 W16630 W16561 H25964 H21754 W01159 W42885 AA176730 H39504 N39788 AA182956 H27585 AA082164 AA328927 AA339934 H61805 H61804 H45580 AA476229 AA714104 AA507471 AI262184 AI139474 AI139476 AI001045 AA614374 AA593153 F33347 F34679 T68225 N25703 AA186999 AI623318 F18313 N72069
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	100771	102/7_1	AW503953 BE466278 AW029058 AI492113 H14384 AA349625 AA324192 AA326695 R52857 AA019306 AA021656 Z42668 AA323883
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<b>50</b> °	123360 116158	333064_1 11903_1	AA532718 AI821485 AI791194 AI821930 AA504784 AW969151 F37127 AA654206 F27974  AA381807 AF144755 NM_013332 AA320807 BE264360 BE312752 AW381329 AW381298 BE301024 AI800437 AI309121 AI343669 AI800457  AA054543 AI310162 AI744870 AI769640 AW674287 AA461187 AW151696 AI277620 AI347821 AA035341 AA670144 AI744879 AI760462  AW514880 AI954915 AI696966 AW105694 AW105695 AI948588 AW083071 BE549300 AW082974 AI862078 AW236191 AW241771 AI368420  AI335595 AI765786 AA054583
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5			Al261532 Al934695 Al923632 Al720333 AW304217 Al089510 Al719846 Al587160 Al498213 AA759077 AW069810 Al304791 Al939998 AA687919 AA852809 T99527 AA527012 AW001996 Al479890 Al863946 Al688036 AA513191 AA863552 N73100 Al368248 AW263462 AA477464 AA617664 AW591651 T29194 Al610373 Al197850 Al364502 Al280917 Al707897 AA290976 D20308 AA285059 Al148183 AA897561 AW408028 AA130549 H75395 AA453330 AA159162 AA158842 BE546820 AA430426 AA428764 AA421577 BE218289 BE540828 BE259495 Al991221 Al800148 BE545935 AA856632 AA402802 AW001345 AA505268 Al831247 AA661521 AW518864 AW193589 Al871010 AA857226 AA758930 AA588803 AA723089 Al719387 AA424374 T95659 AW809033 AW387658 AA761238 AA402306 AA657982 AW609068 AW809046 AW387630 AW387651 AW387554 AA451772 AW609065 AW387608 BE272809 AA464627 AW609555 AW378072 AW579620 AW579594 AA449966
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70			AI5732B2 AW794752 AA370328 AW998896 AW797239 AW998912 AW794742 AI954543 AI810067 AW073373 AA370325 AW195330 C18106 AW998736 R79476 AA429721 AI891081 AI381534 AW022137 AW020000 AI630329 N99428 AI870222 AI971257 AI922196 AI857753 AW579397 D56749 AI925005 AI685727 AW805573 AI982678 AI784604 AI005625 AW877772 AI634947 AI950829 AA493243 BE166086 AI801820 AI925643 AI627992 AW316704 AI261318 D57757 AA887178 AW770406 AI972075 AI222254 AI675794 D58060 AI701954 D58166
			Al799500 AW805669 AW275098 AW874253 Al962991 Al248184 AW996924 Al017462 AW022260 Al885957 BE176841 AA878863 Al97419 AW662094 Al479529 BE177025 D57403 AA507952 AW664593 AW800998 Al985773 AA566089 AA442759 Al624670 Al460284 Al800205
75			AI537788 AI537593 AI244382 AA583463 AA922678 AA864382 AI610837 D58070 AA844283 AA947992 N73801 AI453821 D58184 AI678887 AW243755 AA746085 D57742 AA757380 R44148 AA496403 BE180303 AW363528 BE006616 D57395 AW805507 AW805511 AA617991 AI373585 H30122 D57744 AW805501 D57691 D58148 AW873164 AW768483 D57601 AA777812 AA837997 BE180123 D57599 AA485387 AW022208 D58096 N67917 W95944 AW805506 D57518 D57990 AI074096 D56521 D58161 AA428720 D56648 D57778 AW805504 D57750
80			D58108 AW021706 D57449 D57041 D58277 D56935 Al356974 D57023 AA018712 H27631 D57851 D57514 D57268 D57468 AW805646 Al278945 D57323 D56986 D57539 D57829 D58078 AW805515 Al348684 D57772 R74449 BE041558 D56746 AW798485 D56640 AA985597 D56702 D56849 D56874 AW581419 AA470397 D57591 AW798984 T27640 N66497 D56803 AA618186 AW805647 D57945 N23726 D56637 N23730 D56992 BE176882 BE176839 BE176909 D56757 N68137 D56987 Al559806 AA631437 D57464 D56718 C17030 T29278 D57377

			AW021936 AW118330 AA515358 D56610 AA494092 D56934 T97774 AI473546 R74350 R84834 AA579200 D56616 C03207 D57391 N52416 D56928 R79209 D56925 AA020879 D45546 AI858769 R20750 T09381 F01435 AW627906 D58202 AI933993 F01912 H27552 AA174191 T16515 AW023216 AA434146 H83387 AI346751 V01512 V01512 AA576407 AW365140 AA937471 BE174681 AI568829 AI274663 R85530 AL048225 H33388 AW798734
5	332577	89088_2	AI826268 AW248872 H69511 AI748806 AW779557 AI992254 AI890377 AW151271 AI356374 AI634503 AA777065 AI590131 H37767 AI889058
			H69512 AA046480 N27343 Al573008 AW130925 Al635838 AW594603 AW000790 Al208239 Al275835 AW090294 AA021587 AW273456 AA505726 AW469424 Al400222 Al025723 BE046148 Al128668 BE350462 AW302601 Al299977 AA284809 Al640358 AW470364 Al241794
			AA650048 AW090027 H15377 AW615318 D60021 Al934336 AW118536 Al041281 AA614238 R85918 AW571741 AW516692 AW572232 AW515188 Al798585 Al392825 Z40518 Al869580 AA469975 Al537819 Al810684 Al701744 Al370410 BE383083 Z44676 BE002481 BE002532
10			AA456765 N44196 D60022 C14604 AA021099 AA284872 BE266647 AW249292
	332640	4172_1	BE5684S2 BE297396 AA449593 AW732490 AW069736 BE548667 AA207229 AF044588 NM_003981 BE268994 AW444578 AA471151
			BE250747 AW732555 AA074582 BE336856 AW408764 AA191159 BE092129 AA310614 AW958677 AA312276 AW750027 AW750046
			AW750032 AW750024 AA188893 AW750054 AW408409 AW750030 BE151875 AA478509 N58721 AA195614 H70079 H75580 BE250401 AA454518 AA007263 AA626405 AA417152 AA004230 AA557354 AW863151 AW863181 AA702179 AI924143 AI671185 BE006198 AA190630
15			Al638795 Al609113 Al056239 BE537023 BE464668 AA634413 BE208066 BE208833 AW250803 Al337375 AA478510 BE501624 Al814763
			AW594726 AJ091408 AA827285 AA189108 AW594169 BE618589 BE618040 AL 135398 AA632206 AJ080126 AJ638180 AA725439 AJ379107
			Al288872 H14801 Al679151 Al263619 Al559213 Al679722 W93249 AA552345 AA417030 Al969543 AA534494 Al038181 AA766364 AA573241
			AI754325 AW043937 BE207865 AI291838 N73585 N73539 AW805051 AA808510 AI699813 AW166044 AW104716 H05808 AA248270
20			BE538022 N56013 AA621586 AA149737 D19871 AW192890 N54283 H73339 AA910989 BE273424 BE560082 AW959012 AA313552
20	332732	5436_1	AW750034 BE072537 BE297947 AW732361 AA449336 D29574 AF191019 NM_015516 BE546494 AL110276 R13844 BE313586 BE336912 R18704 R18703 AA045868 T70952 BE336901 T60387 BE149749
	,002702	0400_1	BE271848 BE271902 AA489929 Z45402 T64360 AA305745 AA009451 T95706 H14907 AA299901 C03221 T72431 AW471185 AA335297
			Al269100 AA345072 AW965160 H27581 R48910 H25380 AA335281 AW973283 T79590 AW183447 T64172 AI744097 Al342358 AA336102
25			AA335299 BE208375 A1140834 AA088181 Al860314 Al738613 T70902 R42077 Al884558 AA489798 Al130828 AA009735 H25381 AW612425
25			R48801 H27507 H30105 H44671 Al631362 AA558470 AW014412 AA552059 AA045801 AW589435 Al039657 H14614 AA974256 R42078
			A1245758 T61886 A1559202 A1074139 A1817313 A1041484 AA437138 A1613032 A1147891 A1457945 AW197727 A1074399 A1758636 A1598048
			AA972077 M85390 R36989 R71936 Al867492 T40081 Z41115 AA772775 T41013 Al695691 T40996 Al826822 N93464 AW955524 AA088651

TABLE 1C

Pkey: Unique number corresponding to an Eos probeset

Ref: Sequence source. The 7 digit numbers in this column are Genbank Identifier (Gf) numbers. "Dunham I. et al." refers to the publication entitled "The DNA sequence of human chromosome 22." Dunham I. et al. (1999) Nature 402:489-495.

Strand: Indicates DNA strand from which exons were predicted.

Nt\_position: Indicates nucleotide positions of predicted exons. 5

10	Pkey	Ref	Strand	Nt_position
10	332792	Dunham, I. et.al.	Plus	73381-73768
	333135	Dunham, I. et.al.	Plus	3361208-3361369
	333137	Dunham, I. et.al.	Plus	3367643-3367726
	333138	Dunham, I. et.al.	Plus	3369205-3369323
	333139	Ounham, I. et.al.	Plus	3369495-3369571
15	333516	Dunham, I. et.al.	Plus	5570204-5570390
	333517	Dunham, I. et.al.	Plus	5570729-5570925
	333795	Dunham, I. et.al.	Plus	7807688-7807795
	333796	Dunham, I. et.al.	Plus	7808253-7808319
	333808	Dunham, I. et.al.	Plus	7880600-7880775
20	333809	Dunham, I. et.al.	Plus	7880600-7880775
	333845	Dunham, I. et.al.	Plus	8005832-8005945
	333849	Dunham, I. et.al.	Plus	8018323-8018472
	334101	Dunham, I. et.al.	Plus	9973413-9973550
0.5	334616	Dunham, I. et.al.	Plus	15176123-15176470
25	334891	Dunham, I. et.al.	Plus	19299770-19299944
	334899	Dunham, I. et.al.	Plus	19315168-19315311
	334900	Dunham, I. et.al.	Plus	19315678-19315743
	334902	Dunham, I. et.al.	Plus	19317083-19317195
20	334905	Dunham, I. et.al.	Plus	19322553-19322680
30	334906	Dunham, I. et.al.	Plus	19323493-19323590
	335044	Dunham, I. et.al.	Plus	20842088-20842682
	335149	Dunham, I. et.al.	Plus	21497441-21497587
	335809	Dunham, I. et.al.	Plus	26310772-26310909
25	335810	Dunham, I. et.al.	Plus	26314767-26314849
35	335824	Dunham, I. et.al.	Pius	26376860-26376942
	336054	Dunham, I. et.al.	Plus	29161685-29161937
	336721	Dunham, I. et.al.	Plus	3371522-3371586
	337182	Dunham, I. et.al.	Plus	23934889-23934962
40	337674	Dunham, I. et.al.	Plus	3332616-3332697
40	337675	Dunham, I. et.al.	Plus	3335368-3335505
	337755	Dunham, I. et.al.	Plus	3971764-3971900
	338038	Dunham, I. et.al.	Plus	8138219-8138392
	338316	Dunham, I. et.al.	Plus	17089711-17089988
15	333124	Dunham, I. et.al.	Minus	3318017-3317932
45	333743	Dunham, I. et.al.	Minus	7573218-7573060
	334221	Ounham, I. et.al.	Minus	12730944-12730387
	334222	Dunham, I. et.al.	Minus	12732417-12732289
	334282	Dunham, I. et.al.	Minus	13285293-13285178
50	334502	Dunham, I. et.al.	Minus	14488605-14488526
20	334578	Dunham, I. et.al.	Minus	15004462-15004304
	334951	Dunham, I. et.al.	Minus	20147708-20147502
	335289	Dunham, I. et.al.	Minus	22305950-22305708
	335290	Dunham, I. et.al.	Minus	22309950-22309891
55	335293	Dunham, I. et.al.	Minus	22316408-22316275
55	335682	Dunham, I. et.al.	Minus	25421215-25421093 25761535-25761444
	335753	Dunham, I. et.al.	Minus	
	335755	Dunham, I. et.al.	Minus	25763806-25763747
	335756	Dunham, I. et al.	Minus	25764330-25764251
60	336662	Dunham, I. et.al.	Minus	2158060-2157993
UU	336684	Dunham, I. et.al.	Minus	2158060-2157993
	337603	Dunham, I. et.al.	Minus	1299296-1299194
	338561	Dunham, I. et.al.	Minus	22311966-22311856
	338562	Dunham, I. et.al.	Minus	22312594-22312465
65	339186	Dunham, I. et al.	Minus	32339211-32339097
05	325889	5867087	Plus	223829-223891
	330032	6682596	Plus	85177-85237
	330033	6682596 5867224	Plus	86663-86723
	326213	5867224 6552458	Minus	60751-60927
70	326816	6117842	Plus	198354-198436 94608-94785
70	327110	5867968	Plus	
	327821	5868068	Plus Minus	131060-131232
	328164 328648	6004473	iviinus Plus	27080-27226 424829-424959
	329365	5868838	Minus	424829 <del>-4</del> 24959 107687-107765
75	925000	J. J	IVIII IUD	101001-101100
. —				

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Table 2A lists about 1165 genes selected to have an interesting expression pattern during androgen withdrawal of prostate cancer tissue. These genes were selected by analysis of variance, such that the P value is less than 0.01, the 90th percentile exhibits a minimum of 100 average intensity across all samples, and a comparison of any group means shows a minimum 3 fold change. The interesting expression patterns can be broadly defined into the following categories:

1. Genes that are expressed early in the time course of androgen withdrawal, then drop off in expression, and then express again with emergence of androgen-independence (hi-lo-lo-hi pattern in table 2A).

2. Genes that are expressed early in the time course, then drop off in expression immediately after androgen-withdrawal, and do not express again with emergence of androgenindependence (hi-lo-lo pattern in table 2A).

3. Genes that are expressed early in the time course, then drop off in expression after several days of androgen withdrawal, and do not express again with emergence of androgen-independence (hi-hi-lo-lo pattern in table 2A).

4. Genes that are not expressed early in the time course, but express only with emergence of androgen-independence (lo-to-to-hi pattern in table 2A).

Genes that are not expressed early in the time course, but then express as androgen is withdrawn and continue to express with emergence of androgen-independence (to-lohi-hi pattern in table 2A).

6. Genes that are not expressed early in the time course, but then express as androgen is withdrawn and drop off again with emergence of androgen-independence (to-to-hi-to 15 pattern in table 2A).

Table 28 lists accession numbers for primekeys lacking a unigenelD in table 2A. For each probeset is listed a gene cluster number from which oligonucleotides were designed. Gene clusters were cumpiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

Table 2C lists genomic positioning for primekeys lacking unigene ID's and accession numbers in table 2A. For each predicted exon is listed genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed.

TABLE 2A: ABOUT 1165 GENES SELECTED TO HAVE AN INTERESTING EXPRESSION PATTERN DURING ANDROGEN WITHDRAWAL OF PROSTATE CANCER 25

Pkey: Unique Eos probeset identifier number

ExAccn: Exemplar Accession number, Genbank accession number

UnigenelD: Unigene number 30

Unigene Tille: Unigene gene title Pattern: Broadly defined expression patterns during androgen withdrawal UnicenelD Unicene Title

	Olean-	C A	11-110	11-1	
	Pkey 433412	ExAcon	UnigenelD	Unigene Title	Pattern
35	429097	AV653729	Hs.8185	CGI-44 protein; sulfide dehydrogenase li	lo-lo-hi-lo
55	429097 442731	AK001270	Hs.196086	hypothetical protein FLJ10408	lo-lo-hi-lo
		AI868167	Hs.131044	ESTs	lo-lo-hi-lo
	420820	W26096	Hs.336635	Homo sapiens, clone IMAGE:4179482, mRNA	lo-lo-hi-lo
	422267	AB033044	Hs.114012	KIAA1218 protein	lo-lo-hi-lo
40	416953	N31537	Hs.269046	ESTs	lo-lo-hi-lo
40	413277	H24177	Hs.75262	cathepsin O	lo-lo-hi-lo
	410209	Al583661	Hs.60548	hypothetical prolein PRO1635	lo-lo-hi-lo
	428523	AW974540	Hs.98626	ESTs	lo-lo-hi-lo
	435847	W93821	Hs.39780	CDA017 protein	lo-lo-hi-lo
45	443967	AW294013	Hs.200942	ESTs	lo-lo-hi-lo
43	440838	AA907075	Hs.131307	ESTs	lo-lo-hi-lo
	404054	1100740		Target Exon	lo-lo-hi-lo
	431697	H66740	Hs.38540	ESTs, Weakly similar to ALU4_HUMAN ALU S	lo-lo-hi-lo
	432114	AL036021	Hs.8934	ESTs	lo-lo-hi-lo
50	446397	AW275603	Hs.200712	ESTs	lo-lo-hi-to
30	414094	H15088	Hs.31433	ESTs	lo-lo-hi-to
	424005	AB033041	Hs.137507	vang (van gogh, Drosophila)-like 2	lo-lo-hi-lo
	424401	H67220	Hs.169681	death effector domain-containing	lo-lo-hi-lo
	449749	Al668611	Hs.49760	ESTs	lo-lo-hi-lo
55	458368	BE504731	Hs.138827	ESTs	lo-lo-hi-lo
22	427221	L15409	Hs.174007	von Hippel-Lindau syndrome	lo-lo-hi-lo
	432715	AA247152	Hs.200483	ESTs, Weakly similar to KIAA1074 protein	lo-lo-hi-lo
	425980	AA366951		gb:EST77963 Pancreas tumor III Homo sapi	lo-lo-hi-lo
	412492	AW962604		gb:EST374677 MAGE resequences, MAGG Homo	lo-lo-hi-lo
60	438882	AA827695		gb:od56c02.s1 NCI_CGAP_GCB1 Homo sapiens	lo-lo-hi-to
00	422473	U94780	Hs.117242	meningioma expressed antigen 6 (coiled-c	lo-lo-hi-lo
	404211	41040405	11- 000000	NM_005936:Homo sapiens myeloid/lymphoid	· to-to-hi-to
	423019	AI640185	Hs.283626	ESTs	lo-lo-hi-lo
	443559	AI076765	Hs.269899	ESTs, Moderately similar to ALU8_HUMAN A	lo-lo-hi-lo
65	444291	AI598022	Hs.193989	TAR DNA binding protein	lo-lo-hi-lo
05	428065	A1634046	Hs.157313	ESTs	lo-lo-hi-lo
	442566	R37337	Hs.12111	ESTs	lo-lo-hi-lo
	442202	BE272862	Hs.106534	hypothetical protein FLJ22625	lo-lo-hi-lo
	439456	AI752409	Hs.109314	hypothetical protein FLJ20980	lo-lo-hi-lo
70	423476	AL035633	11- 5044	Human DNA sequence from clone RP5-1046G1	lo-lo-hi-lo
70	437952	D63209	Hs.5944	solute carrier family 11 (proton-coupled	lo-lo-hi-lo
	451987	AA815092	Hs.77554	Homo sapiens cDNA FLJ14967 fis, clone TH	lo-lo-hi-lo
	453408	AI804732	Hs.295963	ESTs	lo-lo-hi-lo
	444004	N39842	Hs.301444	KIAA1673	lo-lo-hi-lo
75	452691	AA164842	Hs.192619	KIAA1600 protein	lo-lo-hi-lo
13	434865	AW050449	Hs.116507	ESTs	io-lo-hi-lo
	440819	AI809444	Hs.202108	ESTs	lo-lo-hi-to
	419526	Al821895	Hs.193481	ESTs	. lo-lo-hi-lo
	422072	AB018255	Hs.111138	KIAA0712 gene product	lo-lo-hi-lo
80	453459	BE047032	Hs.257789	ESTS	lo-lo-hi-lo
30	419038	AW134924	Hs.190325	ESTs	lo-lo-hi-lo
	413243	AA769266	Hs.193657	ESTs	lo-lo-hi-lo
	432079	AW972746		gb:EST384840 MAGE resequences, MAGL Homo	lo-lo-hi-lo

				Par	
	441328	Al982794 R39769	Hs.159473	ESTs ESTs, Moderately similar to ALU8_HUMAN A	lo-lo-hi-lo lo-lo-hi-lo
	416508 451066	A1758660	Hs.206132	ESTs, Moderately Similar to ALOO_HUMAN A	lo-lo-hi-lo
	446017	N98238	Hs.55185	ESTs	lo-lo-hi-lo
5	447104	R19085	Hs.210706	Homo sapiens cDNA FLJ13182 fis, clone NT	lo-lo-hi-lo
	447211	AL161961	Hs.17767	KIAA1554 protein	lo-lo-hi-lo lo-lo-hi-lo
	447765 429540	AW014112 M85776	Hs.161390	ESTs gb:EST02297 Fetal brain, Stratagene (cat	lo-lo-hi-lo
	444314	Al140497		gb:ow76b09.s1 Soares_fetal_liver_spleen_	lo-lo-hi-lo
10	414555	N98569	Hs.76422	phospholipase A2, group IIA (platelets,	lo-lo-hi-lo
	432677	NM_004482		UDP-N-acetyl-alpha-O-galactosamine:polyp	lo-lo-hi-lo
	422091 423028	A1906339 H90946	Hs.97927	ESTs gb;yu86c02.r1 Soares fetal liver spleen	lo-lo-hì-lo lo-lo-hi-lo
	444040	AF204231	Hs.182982	golgin-67	lo-lo-hi-lo
15	441111	A1806867	Hs.126594	ESTs	lo-lo-hi-lo
	418838	AW385224	Hs.35198	ectonucleotide pyrophosphatase/phosphodi	lo-lo-hi-lo
	415999	AA172179	Hs.294029 Hs.211562	ESTS	lo-lo-hi-lo lo-lo-hi-lo
	429615 427774	AF258627 AA278583	Hs.180737	ATP-binding cassette, sub-family A (ABC1 Homo sapiens clone 23664 and 23905 mRNA	lo-lo-hi-lo
20	438585	AA811371	Hs.123362	ESTs	lo-lo-hi-lo
	424776	AI867931	Hs.164595	ESTs	lo-lo-hi-lo
	413786	AW613780	Hs.13500	ESTs	lo-lo-hi-lo
	421077 445837	AK000061 Al261700	Hs.101590 Hs.145544	hypothetical protein ESTs	lo-lo-hi-lo lo-lo-hi-lo
25	449282	AL048056	Hs.23437	Homo sapiens cDNA FLJ13555 fis, clone PL	lo-lo-hi-lo
	414065	AW515373	Hs.271249	Homo sapiens cDNA FLJ13580 fis, clone PL	lo-lo-hi-lo
	432527	AW975028	Hs.102754	ESTs	lo-lo-hi-lo
	412093	BE242691	Hs.14947	ESTs	lo-lo-hi-lo
30	457121 417280	AI743770 AW173116	Hs.180513 Hs.250103	ESTs, Weakly similar to KIAA0822 protein ESTs	lo-lo-hi-lo lo-lo-hi-lo
50	452445	AB002438	Hs.29596	Homo sapiens mRNA from chromosome 5q21-2	lo-lo-hi-lo
	438624	AA889055	Hs.123468	ESTs	lo-lo-hi-lo
	442343	AA992480	Hs.129874	ESTs	lo-lo-hi-lo
25	401416		11 10010	C14000338*:gi[7459502 pir]]S74665 outer	lo-lo-hi-lo
35	437176 451663	AW176909 A1872360	Hs.42346 Hs.209293	calcineurin-binding protein calsarcin-1 ESTs	lo-lo-hi-lo lo-lo-hi-lo
	449295	AW137268	Hs.270954	ESTs	lo-lo-hi-lo
	426848	H72531	Hs.36190	ESTs	to-lo-hi-lo
40	445467	Al239832	Hs.15617	ESTs, Weakly similar to ALU4_HUMAN ALU S	lo-lo-hi-lo
40	418662	Al801098	Hs.151500	ESTs	lo-lo-hi-lo
	416239 428054	AL038450 Al948688	Hs.48948 Hs.266619	ESTs ESTs	lo-lo-hi-lo lo-lo-hi-lo
	435284	AA879470	Hs.96849	Homo sapiens cDNA FLJ11492 fis, clone HE	lo-lo-hi-lo
	424332	AA338919	Hs.101615	ESTs	lo-lo-hi-lo
45	442369	AJ565071	Hs.159983	ESTs	lo-lo-hi-lo
	420717	AA284447	Hs.271887	ESTs	to-to-hi-to
	439584 440260	AA838114 Al972867	Hs.221612 Hs.7130	ESTs copine IV	lo-lo-hi-lo lo-lo-hi-lo
	426269	H15302	Hs.168950	Homo sapiens mRNA; cDNA DKFZp566A1046 (f	lo-lo-hi-lo
50	428398	AI249368	Hs.98558	ESTs	lo-lo-hi-lo
	407276	Al951118	Hs.326736	Homo sapiens breast cancer antigen NY-BR	lo-lo-hi-lo
	409339	AB020686	Hs.54037	ectonucleotide pyrophosphatase/phosphodi	lo-lo-hi-lo
	442150 415787	AJ368158 H01463	Hs.70983 Hs.93534	PTPL1-associated RhoGAP 1 ESTs	lo-lo-hi-lo lo-lo-hi-lo
55	430685	Al690234	Hs.191666	ESTs, Weakly similar to GNMSLL retroviru	lo-lo-hi-lo
	443794	N94104	Hs.29280	ESTs	lo-lo-hi-lo
	446215	AW821329	Hs.14368	SH3 domain binding glutamic acid-rich pr	lo-lo-hi-lo
	441285	NM_002374	Hs.167	microtubule-associated protein 2	lo-lo-hi-lo
60	448738 403746	BE614081		gb:601503815F1 NIH_MGC_71 Horno sapiens c ENSP00000226812*:KIAA1494 protein (Fragm	lo-lo-hi-lo lo-lo-hi-lo
00	434022	R18374	Hs.117956	ESTs	lo-lo-hi-lo
	435714	AA699325	Hs.269880	ESTs	lo-lo-hi-lo
	439848	AW979249		gb:EST391359 MAGE resequences, MAGP Homo	lo-lo-hi-lo
65	421974	AA301270	U- 44000	gb:EST14192 Testis tumor Homo sapiens cD	lo-lo-hi-lo
65	433332 449919	AI367347 AI674685	Hs.44898 Hs.200141	Homo sapiens clone TCCCTA00151 mRNA sequ ESTs	lo-lo-hi-lo lo-lo-hi-lo
	407192	AA609200	115.200141	gb:af12e02.s1 Soares_testis_NHT Homo sap	lo-lo-hi-lo
	436169	AA888311	Hs.17602	Horno sapiens cDNA FLJ12381 fis, clone MA	lo-lo-hi-lo
<b>~</b> 0	418624	AI734080	Hs.104211	ESTs ·	lo-lo-hi-lo
70	432432	AA541323	Hs.115831	ESTs	lo-lo-hi-lo
	426172 401093	AA371307	Hs.125056	ESTs C12000586*:gi]6330167 dbj BAA86477.1  (A	to-lo-hi-lo lo-lo-hi-lo
	426716	NM_006379	Hs.171921	sema domain, immunoglobulin domain (Ig),	lo-lo-hi-lo
	439569	AW602166	Hs.222399	CEGP1 protein	lo-lo-hi-lo
75	451720	AW970985	Hs.290853	ESTs	to-to-hi-lo
	429163	AA884766		gb:am20a10.s1 Soares_NFL_T_GBC_S1 Homo s	lo-lo-hi-lo
	432435	BE218886	Hs.282070	ESTs	lo-lo-hi-lo lo-lo-hi-lo
	408170 433530	AW204516 BE349534	Hs.31835 Hs.281789	ESTs ESTs	10-10-11-10 10-10-hi-10
80	425776	U25128	Hs.159499	parathyroid hormone receptor 2	to-to-hi-to
	430068	AA464964		gb:zx80f10.s1 Soares ovary tumor NbHOT H	lo-lo-ti-lo
	422725	AA315703	Hs.199993	ESTs, Weakly similar to ALUB_HUMAN IIII	lo-lo-hi-lo

					100.00
	432314	AA533447	Hs.312989	ESTs	lo-lo-hi-lo
	434609			gb:yi60c11.r1 Soares placenta Nb2HP Homo	lo-lo-hi-lo
	448760	AA313825	Hs.21941	AD036 protein	lo-lo-tri-lo
_	417381	AF164142	Hs.82042	solute carrier family 23 (nucleobase tra	lo-lo-hi-lo
5	456334	T50392	Hs.271745	ESTs	lo-lo-hi-lo
	435445	AA737345	Hs.294041	ESTs	lo-lo-hi-lo
	411928	AA888624	Hs.197289	rab3 GTPase-activating protein, non-cata	to-lo-hi-lo
	438869	AF075009		gb:Homo sapiens full length insert cDNA	lo-lo-hi-lo
	423932	T95633	Hs.189703	ESTs	to-to-hi-to
10	422222	Al699372	Hs.193247	hypothetical protein DKFZp434A171	lo-lo-hi-lo
	434941	AW073202	Hs.334825	Homo sapiens cDNA FLJ14752 fis, clone NT	lo-lo-hi-lo
	415736		Hs.291872	ESTs	lo-lo-hi-lo
	432722		Hs.326150	ESTs	lo-lo-hi-lo
	435511	AA683336	Hs.189046	ESTs	io-lo-hi-lo
15	432242	AW022715	Hs.162160	ESTs, Weakly similar to ALU4_HUMAN ALU S	lo-lo-hi-lo
	451141	AW772713	Hs.247186	ESTs	lo-lo-hi-lo
	450546	AA010200	Hs.175551	ESTs	lo-lo-hi-lo
	413351	BE086815		ESTs	lo-lo-hi-lo
	439324	AF086134	Hs.94309	ESTs	lo-lo-hi-lo
20	452688	AA721140	Hs.49930	ESTs, Weakly similar to putative p150 (H	
20	415669	NM_005025			lo-lo-hi-lo
	450164	Al239923	Hs.63931	serine (or cysteine) proteinase inhibito ESTs	lo-lo-hi-lo
	417169				lo-lo-hi-lo
		R13550	Hs.246773	ESTs	lo-lo-hi-lo
25	443645	R36475	Hs.24321	Homo sapiens cDNA FLJ12028 fis, clone HE	lo-lo-hi-lo
23	424878	H57111	Hs.221132	ESTs	lo-lo-hi-lo
	449618	A1076459	Hs.15978	KIAA1272 protein	lo-lo-hi-lo
	432572	A1660840	Hs.191202	ESTs, Weakly similar to ALUE_HUMAN !!!!	lo-lo-hi-lo
	400293	N51002	Hs.306480	Homo sapiens mRNA; cDNA DKFZp761E2112 (f	lo-lo-hi-lo
20	431474	AL133990	Hs.190642	CEGP1 protein	lo-lo-hi-lo
30	421674	T10707	Hs.296355	hypothetical protein FLJ23138	la-lo-hi-lo
	438494	AA908678	Hs.130183	ESTs	lo-lo-hi-lo
	425332	AA633306	Hs.127279	ESTs	lo-lo-hì-lo
	451411	AA017492	Hs.135655	EST	lo-lo-hi-lo
2.5	419972	AL041465	Hs.182982	golgin-67	lo-lo-hi-la
35	434804	AA649530	Hs.348148	gb:ns44f05.s1 NCI_CGAP_Alv1 Homo sapiens	lo-lo-hi-lo
	442832	AW206560	Hs.253569	ESTs	lo-lo-hi-lo
	408660	AA525775		ESTs, Moderately similar to PC4259 ferri	lo-lo-hi-lo
	432674	AA641092	Hs.257339	ESTs, Wealty similar to 138022 hypotheti	lo-lo-hi-lo
4.0	448150	A1472167		ESTs	lo-lo-hi-lo
40	450468	AW379075	Hs.141742	Homo sapiens cDNA FLJ12211 fis, clone MA	lo-lo-hi-lo
	452874	AK001061	Hs.30925	hypothetical protein FLJ10199	lo-lo-hi-lo
	412088	A1689496	Hs.108932	ESTs	lo-lo-hi-lo
	443451	Al057404	Hs.58698	ESTs	lo-lo-hi-lo
	453853	AL040600	Hs.188083	ESTs	lo-lo-hi-lo
45	419863	AW952691	Hs.93485	Homo sapiens mRNA; cDNA DKFZp761D191 (fr	lo-lo-hi-lo
	420729	AW964897	Hs.290825	ESTs	lo-lo-hi-lo
	440801	AA906366	Hs.190535	ESTs	lo-lo-hi-to
	407284	AI539227	Hs.214039	hypothetical protein FLJ23556	lo-lo-hi-lo
	428279	AA425310	Hs.155766	ESTs, Weakly similar to A47582 B-cell gr	lo-lo-hi-lo
50	436862	AJ821940		ESTs, Moderately similar to ALU8_HUMAN A	lo-lo-hi-lo
	432340	AA534222		gb:nj21d02.s1 NCI_CGAP_AA1 Homo sapiens	lo-lo-hi-lo
	442048	AA974603		gb:op34f05.s1 Soares_NFL_T_GBC_S1 Homo s	lo-lo-hi-lo
	418781	T41160	Hs.8404	ESTs	lo-lo-hi-lo
_	450642	R39773	Hs.7130	copine IV	lo-lo-hi-lo
55	451661	AB020650	Hs.26777	Homo sapiens, Similar to KIAA0843 protei	lo-lo-hi-lo
	435812	AA700439	Hs.188490	ESTs	lo-lo-hi-lo
	448065	AI459177	Hs.172759	ESTs, Moderately similar to ALU7 HUMAN A	lo-lo-hi-lo
	453486		Hs.173554	ubiquinol-cylochrome c reductase core pr	lo-to-hi-lo
	414312	AA155694	Hs.191060	ESTs	lo-lo-hi-lo
60	438980	AW502384	1.51707000	gb:UI-HF-BR0p-aka-f-12-0-UI.r1 NIH_MGC_5	lo-lo-hi-lo
	408001	AA046458	Hs.95296	ESTs	
	421476	AW953805	Hs.21887	ESTs	lo-lo-hi-lo
	414426	D60745	Hs.25925	Homo sapiens, clone MGC:15393, mRNA, com	lo-lo-hi-lo
	444563	N57057	Hs.284163	ANKHZN protein	lo-lo-hi-lo
65	418771	AA807881	Hs.25329	ESTs .	lo-lo-hi-lo
00	417843	W07361	Hs.22545	Homo sapiens cDNA FLJ12935 fis, clone NT	lo-lo-hi-lo
	415565	AA642449	Hs.48994		lo-lo-hi-lo
	419229	AJ827237	Hs.282884	ESTs, Wealdy similar to AF151800 1 CGI-4 ESTs	lo-lo-hi-lo
	419905				lo-lo-hi-lo
70	452870	AW248229 AW502761	Hs.93659 Hs.30909	protein disulfide isomerase related prot	lo-lo-hi-lo
, 0	449059	AK000566	Hs.98135	KIAA0430 gene product	lo-lo-hi-lo
				hypothetical protein FLJ20559	lo-lo-hi-lo
	416157	NM_003243		transforming growth factor, beta recepto	lo-lo-hi-lo
	439305	AW393883	Hs.98968	hypothetical protein FLJ23058	lo-lo-hi-to
75	419235	AW470411	Hs.288433	neurotrimin	lo-lo-hi-lo
15	416640	BE262478	Hs.79404	neuron-specific protein	lo-lo-hi-lo
	434938	AW500718	Hs.8115	Homo sapiens, clone MGC:16169, mRNA, com	lo-lo-hi-lo
	408177	A1241733	Hs.43871	ESTs	lo-lo-hi-lo
	438459	T49300	Hs.35304	Homo sapiens cDNA FLJ13655 fis, clone PL	lo-lo-hi-lo
80	418381	AA682393	Hs.119237	ESTs	lo-lo-hi-lo
SU	432161	AK000400	Hs.341181	ESTs, Weakly similar to envelope [H.sapi	lo-lo-hi-lo
	418283	S79895	Hs.83942	cathepsin K (pycnodysostosis)	lo-lo-hi-lo
	421443	BE550141	Hs.156148	hypothetical protein FLJ13231	lo-lo-hi-lo

	440040	15010100		h h a come a decembrat	In In Adda
	416619	AF013168	Hs.79393	tuberous sclerosis 1 hypothetical protein FLJ20147	lo-lo-hi-lo lo-lo-hi-lo
	449802 446714	AW901804 W73818	Hs.23984 Hs.110028	ESTs	lo-io-ti-lo
	413195	AA127382	Hs.22404	protease, serine, 12 (neurotrypsin, moto	lo-lo-hi-lo
5	438233	W52448	Hs.56147	ESTs	lo-lo-hi-lo
-	416051	AA835868	Hs.25253	mannosidase, alpha, class 1A, member 1	lo-lo-hi-lo
	438855	AW946276	Hs.6441	Homo sapiens mRNA; cDNA DKFZp586J021 (fr	lo-lo-hi-lo
	425907	AA365752	Hs.155965	ESTs	to-lo-hi-lo
4.0	451295	Al557212	Hs.17132	ESTs, Moderately similar to I54374 gene	lo-lo-hi-lo
10	415443	T07353	Hs.7948	ESTs	lo-lo-hi-lo
	422366	T83882	Hs.97927	ESTs	lo-lo-hi-lo
	435163	AA668884	Hs.19155	ESTs	lo-lo-hi-lo
	426559	AB001914	Hs.170414	paired basic amino acid cleaving system	lo-lo-hi-lo
15	448988	Y09763	Hs.22785	gamma-aminobutyric acid (GABA) A recepto	lo-lo-hi-lo
13	453655	AW960427	Hs.342874	transforming growth factor, beta recepto	lo-lo-hi-lo
	414516 420028	Al307802	Hs.135560	ESTs, Weakly similar to T43458 hypotheti	lo-lo-hi-lo lo-lo-hi-lo
	430223	AB014680 NM_002514	Hs.8786	carbohydrate (N-acetylglucosamine-6-O) s nephroblastoma overexpressed gene	lo-io-hi-lo
	425887	AL049443	Hs.161283	Homo sapiens mRNA; cDNA DKFZp586N2020 (f	lo-lo-hi-lo
20	442577	AA292998	Hs.163900	ESTs	to-to-hi-lo
	424940	AA985308	Hs.283902	ESTs	lo-lo-hi-lo
	428839	AI767756	Hs.82302	Homo sapiens cDNA FLJ14814 fis, clone NT	lo-lo-hi-lo
	443868	W88483	Hs.293650	Homo sapiens mRNA for RGPR-p117, complet	lo-lo-hi-lo
	430334	AI824719	Hs.328700	ESTs	lo-lo-hi-lo
25	439686	W40445	Hs.235857	ESTs, Weakly similar to I38022 hypotheti	lo-lo-hi-lo
	423754	NM_016181		melanoma antigen	to-to-hi-lo
	415205	H71616	Hs.135233	ESTs	lo-lo-hi-lo
	426413	AA377823		gb:EST90805 Synovial sarcoma Homo sapien	lo-lo-hi-lo
20	407204	R41933	Hs.140237	ESTs, Weakly similar to ALU1_HUMAN ALU S	lo-lo-hi-lo
30	430234	N29317	Hs.236463	KIAA1238 protein	lo-lo-hi-lo
	437143	AW204056	Hs.8917	ESTs	lo-lo-hi-hi lo-lo-hi-hi
	445162 415083	AB011131 Al632683	Hs.12376 Hs.27179	piccolo (presynaptic cytomatrix protein) Homo sapiens cDNA FLJ12933 fis, clone NT	lo-lo-hi-hi
	442924	AA533513	Hs.93659	protein disulfide isomerase related prot	lo-lo-hi-hi
35	429536	AA873016	Hs.206097	oncogene TC21	lo-lo-hi-hi
	458584	AF217518	Hs.324136	PTD012 protein	ło-lo-hi-hi
	419647	AA348947	Hs.91816	hypothetical protein	ło-lo-hi-hì
	427201	AB037860	Hs.173933	nuclear factor I/A	lo-lo-hì-hì
4.0	428030	AI915228	Hs.11493	Homo sapiens cDNA FLJ13536 fis, clone PL	lo-lo-hi-hi
40	411779	AA292811	Hs.72050	non-metastatic cells 5, protein expresse	lo-lo-hi-hi
	442482	NM_014039		PTD012 protein	lo-ło-hi-hi
	417458	NM_005655		TGFB inducible early growth response	lo-lo-hi-hi
	438021	AV653790	Hs.324275	WW domain-containing protein 1	lo-lo-hi-hi
45	409799	D11928	Hs.76845	phosphoserine phosphatase-like	lo-lo-hi-hi lo-lo-hi-hi
43	440676	NM_004987		LIM and senescent cell anligen-like doma	lo-lo-hi-hi
	421437 456362	AW821252 AW973003	Hs.104336 Hs.179909	hypothetical protein hypothetical protein FLJ22995	lo-lo-hi-hi
	407686	AW901268	Hs.126043	chromosome 21 open reading frame 51	lo-lo-hi-hi
	431129	AL137751	Hs.263671	Homo sapiens mRNA; cDNA DKFZp434I0812 (f	lo-lo-hi-hi
50	431874	AW610031	Hs.323914	translocase of inner mitochondrial membr	lo-lo-hi-hi
	448072	AI459306	Hs.24908	ESTs	lo-lo-hi-hi
	436860	H12751	Hs.5327	PRO1914 protein	lo-lo-hi-hi
	448770	AA326683	Hs.21992	likely ortholog of mouse variant polyade	lo-lo-hi-hi
<i></i>	428044	AA093322	Hs.301404	RNA binding motif protein 3	lo-lo-hi-hi
55	451468	AW503398	Hs.293663	ESTs, Moderately similar to 138022 hypot	lo-lo-hi-hi
	440278	BE560870	Hs.9052	ESTs, Weakly similar to 2004399A chromos	lo-lo-hi-hi
	441102	AA973905	11- 405700	intermediate filament protein syncoilin	lo-lo-hi-hi
	423942	AF209704	Hs.135723	glycolipid transfer protein	lo-lo-hi-hi lo-lo-hi-hi
60	425254 409324	U91985 W76202	Hs.105658 Hs.343812	DNA fragmentation factor, 45 kD, alpha p lipoic acid synthetase	lo-lo-hi-hi
OO	431707	R21326	Hs.267905	hypothetical protein FLJ10422	lo-lo-hi-hi
	423335	AB018337	Hs.127287	KIAA0794 protein	lo-lo-hi-hi
	429200	AA447871	Hs.194215	ESTs, Weakly similar to 138022 hypotheti	lo-lo-hi-hi
	429898	AW117322	Hs.42366	ESTs	io-lo-hi-hi
65	409604	AW44448	Hs.49124	ESTs	lo-lo-hi-hi
	431797	BE169641	Hs.270134	hypothetical protein FLJ20280	lo-lo-hi-hi
	437576	BE514383		prothymosin, alpha (gene sequence 28)	lo-lo-hi-hi
	415992	C05837	Hs.145807	hypothetical protein FLJ13593	lo-lo-hi-hi
70	458537	W24704	Hs.54773	ESTs	lo-lo-hi-hi
70	417665	AW852858	Hs.22862	ESTs	lo-lo-hi-hi
	422292	AI815733	Hs.114360	transforming growth factor beta-stimulat	lo-lo-hi-hi
	421501	M29971	Hs.1384	O-6-methylguanine-DNA methyltransferase	lo-lo-hi-hi lo-lo-hi-hi
	457952	U25750	Un 16064	Human chromosome 17q21 mRNA clone 1046:1	lo-lo-hi-hi fo-lo-hi-hi
75	414630	8E410857	Hs.16064 Hs.110480	gb:601301177F1 NIH_MGC_21 Homo sapiens c DC12 protein	io-io-ni-ni Io-lo-hi-hi
15	421990 404956	T31811	113.110400	C1003210*:gij6912582 ref NP_036524.1  pe	lo-lo-hi-hi
	436829	AW297958	Hs.163109	ESTs	to-to-hi-hi
	402106	AK002178	. 10. 100 103	hypothetical protein FLJ11316	lo-lo-hi-hi
	404384			NM_020632*:Homo sapiens ATPase, H(+)-tra	lo-lo-hi-hi
80	445123	AI762911	Hs.145369	ESTs	lo-lo-hi-hi
	401757			Target Exon	lo-lo-hi-hi
	439502	AA836672	Hs.130694	ESTs	lo-lo-hi-hi
				1.64	

	400111	A1045700		Eos Control	lo-lo-hi-hi
	405446 401563	AI015709		Homo sapiens mRNA; cDNA DKFZp586l2022 (f C15001262:qil7304981 reflNP_038528.1  ca	ło-lo-hi-hi lo-lo-hi-hi
	402786			C1000887*:gi]12732453[ref]XP_011474.1] C	lo-lo-hi-hi
5	426484	AA379658	Hs.272759	KIAA1457 protein	lo-lo-hi-hi
	414343	AL036166	Hs.323378	coated vesicle membrane protein	lo-lo-hi-hi
	421970	AF227156	Hs.110103	RNA polymerase I transcription factor RR	lo-lo-hi-hi
	422592	BE081857	Hs.94211	rcd1 (required for cell differentiation,	lo-lo-hi-hi
	413431	AW246428	Hs.75355	ubiquitin-conjugating enzyme E2N (homolo	lo-lo-hi-hi
10	426746	J03626	Hs.2057	uridine monophosphate synthetase (orotat	lo-lo-hi-hi
	400237			NM_001087*:Homo sapiens angio-associated	lo-lo-hi-hi
	402532			Target Exon	lo-lo-hi-hi
	402396	AVAPPODDEA	Hs.289292	Target Exon ESTs	lo-lo-hi-hi lo-lo-hi-hi
15	459649 401512	AW298364	NS.209292	NM_014080:Homo sapiens dual oxidase-like	lo-lo-hi-hi
13	448622	AL046508	Hs.270607	ESTs, Weakly similar to STK2_HUMAN SERIN	lo-lo-hi-hi
	400501	74204000	. 13.2.1 0001	ENSP00000251912*:KIAA1617 protein (Fragm	lo-lo-hi-hi
	452324	W81486	Hs.58648	ESTs	lo-lo-hi-hi
	453146	Al338952	Hs.32194	ESTs	lo-lo-hi-hi
20	430445	AW892432	Hs.65307	ESTs	lo-lo-hi-hi
	401750			NM_012448*:Homo sapiens signal transduce	lo-lo-hi-hi
	435236	T03890	Hs.157208	ESTs, Highly similar to ARX MOUSE HOMEOB	lo-lo-hi-hi
	400375	NM_014115		NM_014115*:Homo sapiens PRO0113 protein	lo-lo-hi-hi
25	412151	AA100529	Hs.286232	Homo sapiens cDNA: FLJ23190 fis, clone L	lo-lo-hi-hi
25	410498 405044	AA355749		gb:EST64459 Jurkat T-cells VI Homo sapie NM_014630*:Homo sapiens KIAA0211 gene pr	lo-lo-hi-hi lo-lo-hi-hi
	413169	AW161061	Hs.62954	ESTs. Weakly similar to zinc finger prot	lo-lo-hi-hi
	402101	A44101001	NS.U2334	ENSP0000217725*:Laminin alpha-1 chain p	lo-lo-hi-hi
	455019	AW850818		ab:IL3-CT0220-091199-026-A03 CT0220 Homo	lo-lo-hi-hi
30	446826	AK000626	Hs.16230	hypothetical protein FLJ20619	lo-lo-hi-hi
-	412180	AW898791	Hs.118837	gb:CM0-NN0075-130400-332-f06 NN0075 Homo	lo-lo-hi-hi
	407273	AJ132560		gb:Homo sapiens mRNA for immunoblobulin	lo-lo-hi-hi
	452895	BE389229	Hs.30954	phosphomevalonate kinase	lo-lo-hi-hi
35	416117		Hs.268787	ESTs	lo-lo-hi-hi
33	430934	Al792302 R84694	Hs.248141	potassium inwardly-rectifying channel, s	lo-lo-hi-hi lo-lo-hi-hi
	416309 444578	T80795	Hs.79194 Hs.193702	cAMP responsive element binding protein ESTs	lo-lo-hi-hi
	401966	100733	113.133102	C17000574:gij8923190 ref NP_060178.1  hy	lo-lo-hi-hi
	444850	AW444882	Hs.148483	ESTs	lo-lo-hi-hi
40	403885			Target Exon	lo-lo-hi-hi
	405435			Target Exon	lo-lo-hi-hi
	422694	C06003	Hs.23782	hypothetical protein FLJ12847	lo-lo-hi-hi
	422912	AW405973	Hs.11637	ESTs	lo-lo-hi-hi
45	412748	BE083158	Hs.10862	Homo sapiens cDNA: FLJ23313 fis, clone H	lo-lo-hi-hi
43	403704 440507	H06994		Target Exon gb:yl81b07.r1 Soares Infant brain 1NIB H	lo-lo-hi-hi lo-lo-hi-hi
	405503	1100334		C7000609*:gi 628012 pirt A53933 myosin i	lo-lo-hi-hi
	456123	R00602		gb:ye74c04.r1 Soares fetal liver spleen	lo-lo-hi-hi
	454261	AF216077	Hs.48376	Homo sapiens clone HB-2 mRNA sequence	lo-lo-hi-hi
50	458956	BE220675		gb:ht98f11.x1 NCI_CGAP_Lu24 Homo sapiens	lo-lo-hi-hi
	418367	AA326035	Hs.59236	hypothetical protein DKFZp434L0718	lo-lo-hi-hi
	444553	Al167530	Hs.149380	ESTs	lo-lo-hì-hi
	405811	41400040		NM_024810:Homo sapiens hypothetical prot	lo-lo-hi-hi
55	429461	AI188219	Hs.99311 Hs.164866	ESTs, Weakly similar to HSJ2_HUMAN DNAJ hypothetical protein FLJ22558	lo-lo-hi-hi lo-lo-hi-hi
55	423378 458516	BE313601 BE010749	Hs.255097	ESTs	lo-lo-hi-hi
	404039	DC010743	115.20001	ENSP00000247650*:Hypothetical 177.6 kDa	lo-lo-hi-hi
	454148	AW732837	Hs.42390	nasopharyngeal carcinoma susceptibility	lo-lo-hi-hi
	412678	AA115575	Hs.114914	ESTs	lo-lo-hi-hi
60	449298	AI911333	Hs.171689	ESTs	lo-lo-hi-hi
	405525			NM_002439*:Homo sapiens mutS (E. coli) h	lo-lo-hi-hi
	424576	BE154142	Hs.96833	ESTs	lo-lo-hi-hi
	451601	N92100	Hs.97437 Hs.103931	centrosomal protein 1 DKFZP434B0335 protein	lo-lo-hi-hi lo-lo-hi-hi
65	422395 434333	AA310177 AA186733	Hs.292154	stromal cell protein	lo-lo-hi-hi
05	413509	BE145419		gb:IL5-HT0198-291099-009-E01 HT0198 Homo	lo-lo-hi-hi
	419504	A1088585	Hs.118904	ESTs	lo-lo-hi-hi
	448586	AF285120	Hs.283734	CGI-204 protein	lo-lo-hi-hi
<b>7</b> 0	401209			C12000519:gi 7710046 ref NP_057914.1  ki	lo-lo-hi-hi
70	423554	M90516	Hs.1674	glutamine-fructose-6-phosphate transamin	lo-lo-hi-hi
	439803	AA001021	Hs.6685	thyroid hormone receptor interactor 8	lo-lo-hi-hi
	424593	AA343729	Un 42024	gb:EST49730 Gall bladder I Homo sapiens	lo-lo-hi-hi
	408122	AI432652 NM_001523	Hs.42824 Hs 57697	hypothetical protein FLJ10718 hyaluronan synthase 1	lo-lo-hi-hi lo-lo-hi-hi
75	409958 408214	AL120445	Hs.77823	hypothetical protein FLJ21343	lo-lo-ni-ni lo-lo-hi-hi
, 5	421911	AL041520		gb:DKFZp434G2317_s1 434 (synonym: htes3)	lo-lo-hi-hi
	407813		Hs.40109	KIAA0872 protein	lo-lo-hi-hi
	425211	M18667	Hs.1867	progastricsin (pepsinogen C)	lo-lo-hi-hi
00	442772		Hs.5957	Homo sapiens clone 24416 mRNA sequence	lo-lo-hi-hi
80	419733		Hs.224961	Homo sapiens cDNA FLJ14415 fis, clone HE	lo-lo-hi-hi
	428260	AW290886	Hs.86999	ESTs, Weakly similar to \$65657 alpha-1C- Sec23 (S. cerevisiae) homolog B	lo-lo-hi-hi lo-lo-hi-hi
	427083	NM_006363	173.17.3437	General for reseasoral sustained o	10-10-11I-1II
				165	

	418583	AA604379	Hs.86211	hypothetical protein	lo-lo-hi-hi
	407355	AA846203	Hs.193974	ESTs, Weakly similar to ALU1_HUMAN ALU S	lo-lo-hi-hi
	454003	AA058944	Hs.116602	Homo sapiens, clone IMAGE:4154008, mRNA,	lo-lo-hi-hi
	425322	U63630	Hs.155637	protein kinase, DNA-activated, catalytic	lo-lo-hi-hi
5	402240	-		Target Exon	lo-lo-hi-hi
_	421867	AA481078	Hs.109045	hypothetical protein FLJ10498	lo-lo-hi-hi
	408603	R25283	Hs.326416	Homo sapiens mRNA; cDNA DKFZo564H1916 (f	lo-lo-hi-hi
	437389				
		AL359587	Hs.271586	hypothetical protein DKFZp762M115	lo-lo-hi-hi
10	457148	AF091035	Hs.184627	KIAA0118 protein	lo-lo-hi-hi
10	400277			Eos Control	lo-lo-hi-hi
	400995			C11000295*:gi[12737279 ref XP_012163.1]	lo-lo-hi-hi
	400818			Target Exon	lo-lo-hi-hi
	402758			C1001899*:gi 12722636 ref XP_010672.1] e	lo-lo-hi-hi
	403708			Target Exon	lo-lo-hi-hi
15	405610			ENSP00000241065*:CDNA	lo-to-hi-hi
15	414242	AA749230	Hs.26433	dolichyl-phosphate (UDP-N-acetylglucosam	lo-lo-hi-hi
		X78592	Hs.99915		
	420757	A/0092	HS.99915	androgen receptor (dihydrotestosterone r	lo-lo-hi-hi
	400965			C11002190*:gi 12737279 ref XP_012163.1	lo-lo-hi-hi
20	401192			Target Exon	lo-lo-hi-hi
20	404407			Target Exon	lo-lo-hi-hi
	401405			Target Exon	lo-lo-hi-hi
	403055			C2002219*:gij12737280jrefjXP_006682.2j k	lo-lo-hi-hi
	404661			C9000306*:gi[12737280]ref[XP_006682.2] k	lo-lo-hi-hi
	433627	AF078866	Hs.284296	Homo sapiens cDNA: FLJ22993 fis, clone K	lo-lo-hi-hi
25	410204	AJ243425	Hs.326035	early growth response 1	lo-lo-hi-hi
20	432642	BE297635	Hs.3069		
		BE29/035	HS.3009	heat shock 70kD protein 9B (mortalin-2)	lo-lo-hi-hi
	400769			Target Exon	lo-lo-hi-hi
	433980	AA137152	Hs.286049	phosphoserine aminotransferase	lo-lo-hi-hi
~ ~	403725			Target Exon	lo-lo-hi-hi
30	413587	AA156164	Hs.286241	protein kinase, cAMP-dependent, regulato	lo-lo-hi-hi
	422614	AI908006	Hs.295362	Homo sapiens cDNA FLJ14459 fis, clone HE	to-to-hi-hi
	400275			NM_006513*:Homo sapiens seryl-IRNA synth	lo-to-hi-hi
	402810			NM_004930*:Homo sapiens capping protein	lo-lo-hi-hi
	452049	BE268289	Hs.27693	peptidylprolyl isomerase (cyclophilin)-l	lo-lo-hi-hi
35	445677	H96577	Hs.6838		
55				ras homolog gene family, member E	lo-lo-hi-hi
	428770	AK001667	Hs.193128	hypothetical protein FLJ10805	lo-lo-hi-hi
	428403	Al393048	Hs.326159	leucine rich repeat (in FUI) interactin	lo-lo-hi-hi
	434647	W74158	Hs.103189	lipopolysaccharide specific response-68	lo-lo-hi-hi
40	402807			ENSP00000235229:SEMB.	lo-lo-hi-hi
40	413992	W26276	Hs.136075	RNA, U2 small nuclear	lo-lo-hi-hi
	407191	AA608751		gb:ae56h07.s1 Stratagene lung carcinoma	to-to-hi-to
	403328			Target Exon	lo-lo-hi-hi
	411984	NM_005419	Hs.72988	signal transducer and activator of trans	lo-lo-hi-lo
	451017	BE391847	Hs.181173	hypothetical protein MGC10771	lo-lo-hi-hi
45	404108	bedo io ii	***************************************	C7000911*:gi 4235142 gb AAD14470.1  (AC0	lo-lo-hi-hi
1.5	407819	R42185	Hs.102720	ESTs	lo-lo-hi-hi
	435876	AW612586	Hs.160271		
			ns.1002/1	G protein-coupled receptor 48	lo-lo-hi-lo
	436716	Al433540		gb:ti69g05.x1 NCi_CGAP_Kid11 Homo sapien	lo-lo-hi-hi
50	401419			Target Exon	lo-lo-hi-hi
50	424363	AW512144	Hs.346947	ESTs, Weakly similar to A48809 carboxyle	lo-lo-hi-hi
	408866	AW292096	Hs.255036	ESTs	lo-lo-hi-hi
	415516	F11411		gb:HSC2WF081 normalized infant brain cDN	lo-lo-hi-hi
	423144	AW851527	Hs.253677	ESTs, Weakly similar to I38022 hypotheti	lo-lo-hi-hi
_	452560	BE077084	Hs.99969	ESTs	lo-lo-hi-hi
55	439827	AA846538	Hs.187389	ESTs	lo-lo-hi-hi
	419709	AA255592	Hs.347973	ESTs, Weakly similar to alternatively sp	lo-lo-hi-hi
	413672		113.541313		
		BE156536 AA354572		gb:QV0-HT0368-310100-091-h10 HT0368 Homo	lo-lo-hi-hi
	425291		Lle 0574 40	gb:EST62857 Jurkat T-cells V Horno sapien	lo-lo-hi-hi
60	427403	AA402107	Hs.257146	ESTs, Moderately similar to I38022 hypot	lo-lo-hì-hi
60	430911	AW937461	Hs.255377	ESTs	lo-lo-hi-hi
	435293	A1040777	Hs.117170	ESTs	lo-lo-hi-hi
	448490	AI523897	Hs.271692	ESTs, Weakly similar to I38022 hypotheti	lo-lo-hi-hi
	449539	W80363	Hs.58446	ESTs	lo-lo-hi-hi
	458082	AW978811	Hs.314451	ESTs, Weakly similar to ALU1_HUMAN ALU S	lo-lo-hi-hi
65	459407	N92114		gb:za22h11.r1 Soares fetal liver spleen	lo-lo-hi-hi
05	423231	AA323486	Hs.271273	Homo sapiens cDNA FLJ12335 fis, clone MA	lo-lo-hi-hi
		AW382884			
	450628		Hs.204715 Hs.136075	ESTs	lo-lo-hi-hi
	411690	AA669253		RNA, U2 small nuclear	lo-lo-hi-hi
70	414739	U83867	Hs.77196	spectrin, alpha, non-erythrocytic 1 (alp	lo-lo-hi-hi
70	444169	AV648170	Hs.58756	ESTs	to-lo-hi-hi
	420911	U77413	Hs.100293	O-linked N-acetylglucosamine (GlcNAc) tr	lo-lo-hi-hi
	422195	AB007903	Hs.113082	KIAA0443 gene product	lo-lo-hi-hi
	452704	AA027823	Hs.149424	Homo sapiens PNAS-130 mRNA, complete cds	lo-lo-hi-hi
	425074	AA495930		Homo sapiens cDNA: FLJ22165 fis, clone H	lo-lo-hi-hi
75	426376	N46752	Hs.302985	ESTs	lo-lo-hi-hi
. •	447754	AW073310	Hs.163533	Homo sapiens cDNA FLJ14142 fis, clone MA	lo-lo-hi-hi
		AI469213	Hs.71404		
	413686			ESTs	lo-lo-hi-hi
	449000	U69560	Hs.3826	kelch-like protein C3IP1	io-lo-hi-hi
00	430064	AK000091	Hs.231436	hypothetical protein FLJ20084	lo-lo-hi-hi
80	412205	N33818	Hs.20274	ESTs, Weakly similar to unnamed protein	lo-lo-hì-hì
	423955	AJ420582	Hs.136164	cutaneous T-cell lymphoma-associated turn	lo-lo-hì-hi
	455619	8E063853		gb:QV3-BT0296-011299-022-g09 BT0296 Homo	lo-lo-hi-hi
				166	

	•				
	40872				lo-lo-hi-h
	45971 41791				lo-lo-hi-h
_	40296	4		NM_022095*:Homo sapiens hypothetical C2H	lo-lo-hi-hi lo-lo-hi-hi
5	42438			3 ANKHZN protein	lo-lo-hi-hi
	42722 41045			3 RNA binding motif protein 6 gb:RC3-BT0316-270400-016-f10 BT0316 Homo	lo-lo-hi-hi
	40071	3	•	NM_006165*:Homo sapiens nuclear factor r	lo-lo-hi-hi lo-lo-hi-hi
10	40721			ubiquilin-conjugating enzyme E2H (homolo	lo-lo-hi-hi
10	44931: 41961:		Hs.223666 Hs.110613		ło-lo-hi-hi
	455272			gb:RC4-HT0231-041199-012-b04 HT0231 Homo	lo-lo-hi-hi lo-lo-hi-hi
	401839	_		NM_005177*:Homo sapiens ATPase, H+ trans	lo-lo-hi-hi
15	440422 436819			myosin phosphatase, target subunit 2	lo-lo-hi-hi
	413644				lo-lo-hi-hi
	413939			ESTs, Weakly similar to ALU7 HUMAN ALU S	lo-lo-hi-hi lo-lo-hi-hi
	448198 450488				lo-lo-hi-hi
20	433507			ESTs, Moderately similar to HPV16 E1 pro ESTs	lo-lo-hi-hi
	438996		6 Hs.110613	KIAA0421 protein	io-lo-hi-lo Io-lo-hi-lo
	442789 407251		1 Hs.131191		lo-lo-hi-lo
	409051		·	transaldolase 1 gb:zn04d03.r1 Stratagene hNT neuron (937	lo-lo-hi-lo
25	409123	AA063403		gb:zm04d12.s1 Stratagene corneal stroma	lo-lo-hi-lo lo-lo-hi-lo
	416225 433735			ESTs, Weakly similar to PC4259 ferritin	lo-lo-hi-to
	434404				lo-lo-hi-lo
20	446667	BE161878			lo-lo-hi-lo lo-lo-hi-lo
30	447982 438890		Hs.137551		lo-lo-hi-lo
	427882				lo-lo-hi-lo
	459680	H96982	Hs.42321	ESTs	lo-lo-hi-lo lo-lo-hi-lo
35	416632 453876		Hs.141304	ESTs	lo-lo-hi-lo
22	414528	AA148950		ESTs, Wealdy similar to 138022 hypotheti ESTs	lo-lo-hi-lo
	419902	AA804409	Hs.118920	ESTs	lo-lo-hi-lo lo-lo-hi-lo
	409542 433560	AA503020	Hs.36563	hypothetical protein FLJ22418	lo-lo-hi-lo
40	447499	AI925195 AW262580	Hs.130891 Hs.147674	hypothetical protein MGC4400 protocadherin beta 16	lo-lo-hi-lo
	435023	Al692552	110.171017	gb:wd73f12.x1 NCI_CGAP_Lu24 Homo saciens	lo-lo-hi-lo lo-lo-hi-lo
	412156	H29487	Hs.17110	Homo sapiens mRNA; cDNA DKFZp434C2016 (f	lo-lo-hi-lo
	414505 404277	R45389	Hs.23558	ESTs, Weakly similar to A48042 lysosomal NM_019111*:Homo sapiens major histocompa	lo-lo-hi-lo
45	414662	AL036058	Hs.76807	major histocompalibility complex, class	lo-lo-hi-lo lo-lo-hi-lo
	444430	Al611153	Hs.6093	Homo sapiens cDNA: FLJ22783 fis, clone K	lo-lo-hi-lo
	445612 403739	N94126	Hs.12969	hypothetical protein ENSP00000251563*:UDP-glucuronosyltransfe	lo-lo-hi-lo
50	403740			NM_001076*:Homo sapiens UDP glycosyltran	lo-lo-hi-lo lo-lo-hi-lo
50	411084	T18987	Hs.125472	ESTs, Moderately similar to KIAA0877 pro	lo-lo-hi-lo
	429143 443060	AA333327 D78874	Hs.197335 Hs.8944	plasma glutamate carboxypeptidase procollagen C-endopeptidase enhancer 2	lo-lo-hi-lo
	422749	W01076	Hs.278573	CD59 antigen p18-20 (antigen identified	lo-lo-hi-lo lo-lo-hi-lo
55	429441	AJ224172	Hs.204096	lipophilin B (uteroglobin family member)	lo-lo-hi-lo
55	414382 441560	AW380339 F13386	Hs.8068 Hs.7888	hematopoietic PBX-interacting protein Homo sapiens clone 23736 mRNA sequence	lo-lo-hi-lo
	446106	AA377165	Hs.44833	ESTs	lo-lo-hi-lo lo-lo-hi-lo
	452239	AW379378	Hs.170121	protein tyrosine phosphatase, receptor t	lo-lo-hi-lo
60	446874 412795	AW968304 BE241753	Hs.56156 Hs.74592	ESTs special AT-rich sequence binding protein	lo-lo-hi-lo
	430325	AF004562	Hs.239356	syntaxin binding protein 1	lo-lo-hi-lo lo-lo-hi-lo
	426392	AW968324	Hs.17384	ESTs	lo-lo-hi-lo
	447448 415743	BE244285 AA167664	Hs.14333	F-box only protein 29 ESTs, Weakly similar to Z195_HUMAN ZINC	lo-lo-hi-lo
65	431607	AB033097	Hs.183669	KIAA1271 protein	lo-lo-hi-lo lo-lo-hi-lo
	411979	X85134	Hs.72984	retinoblastoma-binding protein 5	lo-lo-hi-lo
	453620 431099	BE396163 Y13367	Hs.25005 Hs.249235	ESTs, Weakly similar to ALU5_HUMAN ALU S phosphoinositide-3-kinase, class 2, alph	lo-lo-hi-lo
70	421687	AL035306	Hs.106823	hypothetical protein MGC14797	lo-lo-hi-lo
70	439565	AF086386	Hs.145599	ESTs	lo-lo-tri-lo bo-lo-tri-lo
	442349 410096	W40516 AW245200	Hs.132355 Hs.267400	Homo sapiens cDNA: FLJ22119 fis, clone H	lo-lo-hi-lo
	429447	AW812452	Hs.83286	hypothetical protein MGC5540 ESTs, Weakly similar to S14747 sphingomy	lo-lo-hi-lo
75	431802	AL133570	Hs.270571	Homo sapiens mRNA; cDNA DKFZp434L201 (fr	lo-lo-hi-lo lo-lo-hi-lo
75	441715 458230	Al929453 BE311851	Hs.342655	Homo sapiens cDNA FLJ13289 fis, clone OV	ìo-lo-hi-lo
	428788	AF082283	Hs.6639 Hs.193516	KIAA1624 protein B-cell CLL/lymphoma 10	lo-lo-hi-lo
	450818	Al740573	Hs.142827	P311 protein	lo-lo-hi-lo lo-lo-hi-lo
80		AK002060 AE150003	Hs.91251	hypothetical protein FLJ11198	lo-lo-hi-lo
00		AF159093 Al921573	Hs.213107	Homo sapiens endogenous retrovirus RAN1 ESTs	lo-lo-hi-lo
		AA046772		RNA binding motif protein, X chromosome	lo-lo-hi-lo lo-lo-hi-lo
					- TOTIFO

	423749	U09848	Hs.132390	zinc finger protein 36 (KOX 18)	lo-lo-hi-lo
	428898	AB033070	Hs.194408	KIAA1244 protein	lo-lo-hi-lo
	458258	AW406546	Hs.127971	ESTs .	lo-lo-hi-lo
_	429521	BE048708	Hs.50949	ESTs	lo-lo-hi-lo
5	402185			Target Exon	lo-io-hi-lo
	415961	H10983	Hs.155919	ESTs	lo-lo-hi-lo
	457265	AB023212	Hs.225967	KIAA0995 protein	lo-lo-hi-lo
	412419	AW948630		gb:QV0-FT0001-050500-226-g05 FT0001 Homo	lo-lo-hi-lo
10	438397		, Hs.123206	ESTS	lo-lo-hi-lo
10	440509	BE410132	Hs.134202	ESTs, Weakly similar to T17279 hypotheti	lo-lo-hi-lo
	423895	AA332215		gb:EST36124 Embryo, 8 week I Homo sapien	lo-lo-hi-lo
	400251	AMMOCACO	U- 447900	NM_004651*:Homo sapiens ubiquitin specif	lo-lo-hi-lo lo-lo-hi-lo
	445094	AW296163	Hs.147296	ESTS	lo-lo-hi-lo
15	432323 444290	AK001409 AA262496	Hs.274356	hypothetical protein FLJ10547 gb:zs20f11.r1 NCI_CGAP_GCB1 Homo sapiens	lo-lo-hi-lo
13	435803	Z44194	Hs.4994	transducer of ERBB2, 2	lo-lo-hi-lo
	436905	N31273	Hs.42380	ESTs	lo-lo-hi-lo
	401849	110.270	110,42000	Targel Exon	lo-lo-hi-lo
	402249			C19000553*:gi]12741444]ref[XP_008888.2]	lo-lo-hi-lo
20	406180	AB018249		small inducible cytokine subfamily A (Cy	lo-lo-hi-lo
	448176	AI672546	Hs.170507	ESTs	lo-lo-hi-lo
	409259	AW608930	Hs.52184	hypothetical protein FLJ20618	lo-lo-hi-lo
	457335	AW969834	Hs.303303	ESTs	lo-lo-hi-lo
	452444	BE144022		gb:MR0-HT0165-191199-004-f05 HT0165 Homo	lo-lo-hi-lo
25	405429			Target Exon	lo-lo-hi-lo
	430103	AA465259		gb:aa33b03.r1 NCI_CGAP_GCB1 Homo sapiens	lo-lo-hi-lo
	439944	AA856767	Hs.124623	ESTs	lo-lo-hi-lo
	411283	AW852754		gb:PM1-CT0247-180100-009-c05 CT0247 Homo	lo-lo-hi-lo
~ ~	458195	R10085	Hs.130370	ESTs	lo-lo-hi-lo
30	452654	BE004783		gb:MR2-BN0114-270400-004-e11 BN0114 Homo	lo-lo-hi-lo
	425684	AF000989	Hs.159201	thymosin, beta 4, Y chromosome	lo-lo-hi-lo
	429452	Al949495	Hs.133998	Homo sapiens cDNA FLJ13202 fis, clone NT	lo-lo-hi-lo
	431709	AF220185	Hs.267923	uncharacterized hypothalamus protein HTO	lo-lo-hi-lo
35	411701	BE181659	11- 204202	gb:QV1-HT0638-070500-191-g07 HT0638 Homo	lo-lo-hi-lo
55	430729 447476	AJ572560	Hs.301283 Hs.20880	KIAA0793 gene product ESTs, Weakly similar to I38022 hypotheti	lo-lo-hi-lo lo-lo-hi-lo
	450436	BE293466 AW293661	Hs.131887	ESTs	lo-lo-hi-lo
	405365	A11233001	115.151007	CX001212*:gi[7861932]gb[AAF70445.1] (AF2	lo-lo-hi-lo
	419555	AA244416		gb:nc07d11.s1 NCI_CGAP_Pr1 Homo sapiens	lo-lo-hi-lo
40	446103	U90918	Hs.13804	hypothetical protein dJ462O23.2	lo-lo-hi-lo
	400986			NM_024085*:Homo sapiens hypothetical pro	lo-lo-hi-lo
	424194	BE245833	Hs.169854	gb:TCBAP1E1908 Pediatric pre-B cell acut	lo-lo-hi-lo
	400210			Eos Control	lo-lo-hi-lo
	400234			NM_005336:Homo sapiens high density lipo	lo-lo-hi-lo
45	400235			NM_005336:Homo sapiens high density lipo	lo-lo-hi-lo
	405387			NM_022170*:Homo sapiens Williams-Beuren	lo-lo-hi-lo
	433075	NM_002959		sortilin 1	lo-lo-hi-lo
	406302			C16000922:gi]7499103 pir [T20903 hypothe	lo-lo-hi-to
50	428181	AA423976		gb:zv62h06.s1 Soares_testis_NHT Homo sap	lo-lo-hi-lo
50	456629	AW891965	Hs.279789	histone deacetylase 3	lo-lo-hi-lo
	426940	AA393537	Hs.98347	ESTs, Weakly similar to JC5308 testis-sp	lo-lo-hi-lo
	433555	AA535902	Hs.146211	Homo sapiens HERC2P7 pseudogene, partial ESTs	lo-io-hi-io lo-io-hi-io
	421431	AA650117 Al554923	Hs.283107	gb:le53h12.x1 Soares_NFL_T_GBC_S1 Homo s	lo-lo-hi-lo
55	448631 433521	T66087	Hs.112482	Homo sapiens unknown mRNA sequence	to-to-ti-to
55	407187	AA446971	110.112402	gb:zw85f11.s1 Soares_total_fetus_Nb2HF8_	lo-lo-hi-lo
	450739	AI732707	Hs.116506	ESTs, Weakly similar to ALU7_HUMAN ALU S	lo-lo-hi-lo
	440004	BE397117	Hs.120824	hypothetical protein FLJ21845	lo-lo-hi-lo
	403947	NM_005032		plastin 3 (T isoform)	lo-lo-hi-lo
60	405529	AW410458		chromosome 11 open reading frame2	lo-lo-hi-lo
	402163			C19001075*:gij4567179jgbJAAD23607.1JAC00	lo-lo-hi-lo
	404663			ENSP00000251884:KIAA1521 protein (Fragme	lo-lo-hi-lo
	400220			Eos Control	lo-lo-hi-lo
~~	401444			Target Exon	lo-lo-hi-lo
65	455824	BE143703		gb:MR0-HT0164-191199-004-f03 HT0164 Homo	lo-lo-hi-lo
	400206			Eos Control	lo-lo-hi-lo
	458659	AW749895	Hs.332520	Homo sapiens mRNA; cDNA DKFZp434A1014 (f	lo-lo-hi-lo
	428666	AL080190	Hs.189242	Homo sapiens mRNA; cDNA DKFZp434A202 (fr	lo-lo-hi-lo
70	428442		Hs.98606	ESTS	lo-lo-hi-lo
70	440151	AA868167	No 240426	gb:ak38e07.s1 Soares_testis_NHT Homo sap	lo-lo-hì-lo lo-lo-hi-lo
	431046 443914	AW854382 Al091173	Hs.249126 Hs.222362	Homo sapiens clone 24894 mRNA sequence ESTs, Weakly similar to p40 [H.sapiens]	10-10-111-10 10-10-111-10
	402469	M031173	1 13.222302	Target Exon	to-to-hi-to
	418155	R45481	Hs.23719	ESTs, Weakly similar to I38022 hypotheti	lo-lo-hi-lo
75	446893	Al610818	Hs.7110	ESTs	to-to-hi-to
. •	442336	AW340958	Hs.7572	ESTs	lo-lo-hi-lo
	421290	NM_014368		LIM homeobox protein 6	lo-lo-hi-lo
	450374	AA397540	Hs.60293	Homo sapiens clone 122482 unknown mRNA	lo-lo-hi-lo
0.0	402347			Target Exon	lo-lo-hi-lo
80	415184	AA380436	Hs.211973	homolog of Yeast RRP4 (ribosomal RNA pro	lo-lo-hi-lo
	415632	U67085	Hs.78524	TcD37 homolog	lo-lo-hi-lo
	423718	AL119520	Hs.180737	Homo sapiens clone 23664 and 23905 mRNA	lo-lo-hi-lo

	440440	A1A/042040	Lie ananna	ECT.	1. 1. 1.1.
	449140 431241	AW013840 AA496799	Hs.202092 Hs.36958	ESTs ESTs	lo-lo-hi-lo
	416631	H69466	NS.30330	gb:yr88f07.r1 Soares fetal liver spleen	lo-lo-hi-lo lo-lo-hi-lo
	424168		Hs.321677	signal transducer and activator of trans	lo-lo-hi-lo
5	401600	BE247275	113.321077	U5 snRNP-specific protein, 116 kD	lo-lo-hi-lo
•	420588	AF000982	Hs.147916	DEAD/H (Asp-Glu-Ala-Asp/His) box polypep	lo-lo-hi-lo
	414111	BE047679	Hs.152982	hypothetical protein FLJ13117	to-lo-hi-lo
	417138	AA193646	Hs.65771	Homo sapiens chromosome 19, BAC CIT-HSPC	lo-lo-hi-lo
10	424318	AA476515	Hs.172723	ESTs	lo-lo-hi-lo
10	455653	BE154075		gb:PM0-HT0339-200400-010-E05 HT0339 Homo	lo-lo-hi-to
	451493	H38656	Hs.32854	ESTs	lo-lo-hi-lo
	457015	AA688058	Hs.261544	ESTs	lo-lo-hi-lo
	403654	*******	11.004007	NM_003071:Homo sapiens SWI/SNF related,	lo-lo-hi-lo
15	435203	AW957127	Hs.294027	ESTs	lo-lo-hi-lo
13	409322 437764	BE091159 AA767795	Hs.22687 Hs.166832	ESTs, Moderately similar to unnamed prot ESTs	lo-lo-hi-lo
	432542	AW083920	Hs.16098	claudin 2	lo-lo-hi-lo lo-lo-hi-lo
	436125	AA765895	Hs.152895	ESTs	lo-lo-hi-lo
	403217	AL134878	110.102000	ribosomal protein, large P2	lo-lo-hi-lo
20	434023	Al277883	Hs.146141	ESTs	lo-lo-hi-lo
	442419	AI749893	Hs.270532	ESTs, Weakly similar to 138022 hypotheti	lo-lo-hì-lo
	443667	Al129066	Hs.135457	ESTs	lo-lo-hi-lo
	451445	AA017609	Hs.343449	gb:ze37e01.r1 Soares retina N2b4HR Homo	lo-lo-hi-lo
0.5	454775	BE160229		gb:QV1-HT0413-090200-062-a12 HT0413 Homo	lo-lo-hi-lo
25	411053	AW815061		gb:CM0-ST0209-271099-082-d10 ST0209 Homo	lo-lo-hi-lo
-	435312	AJ243396	Hs.4865	voltage-gated sodium channel beta-3 subu	lo-lo-hi-lo
	450875	AK000724	Hs.301553	karyopherin alpha 6 (importin alpha 7)	lo-lo-hi-lo
	451180	H61899	Hs.171937	steroid dehydrogenase-like .	lo-lo-hi-lo
30	427327 444321	AW501456 AW204210	Hs.288283	Homo sapiens cDNA: FLJ22355 fis, clone H	lo-lo-hi-lo
50	405109	N47812	Hs.122275	Homo sapiens mRNA; cDNA DKFZp564N1623 (f CGI-35 protein	lo-lo-hi-lo
	450182	A1796400	Hs.240767	Human DNA sequence from clone RP1-12G14	lo-lo-hi-lo lo-lo-hi-lo
	424990	AU076896	Hs.154095	zinc finger protein 143 (clone pHZ-1)	lo-lo-hi-lo
	428997	AF065391	Hs.194718	zinc finger protein 265	lo-lo-hi-lo
<b>35</b> .	402602			NM_021186*:Homo sapiens zona pellucida g	lo-to-hi-lo
	428772	Al524039	Hs.192524	ESTs	lo-lo-hi-lo
	423759	Al142358	Hs.184361	ESTs, Moderately similar to ALU7_HUMAN A	lo-lo-hi-lo
	434350	AL042940	Hs.93872	KIAA1682 protein	lo-lo-hi-lo
40	442274	A)733484	Hs.129182	ESTs	lo-lo-hi-lo
40	442884	A1076570	Hs.134053	ESTs	lo-lo-hi-lo
	400481	TE4000		Target Exon	lo-lo-hi-lo
	407283	T51008	U= 250004	gb:yb55e08.s1 Stratagene ovary (937217)	lo-lo-hi-lo
	408859 455615	AW291672 BE045344	Hs.258981 Hs.274923	ESTs ESTs, Moderately similar to unnamed prot	lo-lo-hi-lo
45	427315	AA179949	Hs.175563	Homo sapiens mRNA; cDNA DKFZp564N0763 (f	lo-lo-hi-lo lo-lo-hi-lo
	449375	R07114	Hs.271224	ESTs	lo-lo-hi-lo
	419937	AB040959	Hs.93836	DKFZP434N014 protein	lo-lo-hi-lo
	422231	AA443512	Hs.101383	ESTs	lo-lo-hi-lo
50	437210	AA311443	Hs.293563	Homo sapiens mRNA; cDNA DKFZp586E2317 (f	lo-lo-hi-lo
50	418056	AA524886		gb:nh34f02.s1 NCI_CGAP_Pr3 Homo sapiens	lo-lo-hi-lo
	446586	N58790	Hs.268820	ESTs	lo-lo-hi-lo
	407949	W21874	Hs.247057	ESTs, Weakly similar to 2109260A B cell	lo-lo-hi-lo
	440296	D30829	Hs.180610	splicing factor proline/glutamine rich (	lo-lo-hi-lo
55	422260 434685	AA315993 AA642445	Hs.105484 Hs.287467	regenerating gene type IV Homo sapiens cDNA FLJ11948 fis, clone HE	lo-lo-hi-lo
55	412657	AW976165	NS.201401	gb:EST388274 MAGE resequences, MAGN Homo	lo-lo-hi-lo
	405188	A11310103		Target Exon	io-lo-hi-lo lo-lo-hi-lo
	416954	Al222358		gb:qh04c12.x1 Soares_NFL_T_GBC_S1 Homo s	lo-lo-hi-lo
	423700	AA232375	Hs.58606	SNRPN upstream reading frame	lo-lo-hi-lo
60	430288	BE394943	Hs.13804	hypothetical protein dJ462O23.2	lo-lo-hi-lo
	435184	T67162	Hs.135127	ESTs, Weakly similar to unnamed protein	lo-lo-hi-lo
	431475	A1567669	Hs.40342	putative nuclear protein	lo-lo-hi-lo
	445239	AI217375	Hs.170023	ESTs, Weakly similar to CA36_HUMAN COLLA	lo-lo-hi-lo
65	436151	AK000801	Hs.324271	Homo sapiens cDNA FLJ20794 fis, clone CO	lo-lo-hi-lo
03	448489	A1523875	11- 000500	gb:tg97d04.x1 NCI_CGAP_CLL1 Homo sapiens	lo-lo-hi-lo
	424470	BE244261	Hs.323502	Homo sapiens cDNA: FLJ23539 fis, clone L ESTs	lo-lo-hi-lo
	434733 409469	Al334367 AW517236	Hs.159337 Hs.335762	ESTS	lo-lo-hi-lo
	414034	U89277	Hs.305985	early development regulator 1 (homolog o	lo-lo-hi-lo
70	420382	AW959165	Hs.270034	Homo sapiens, Similar to nuclear localiz	lo-lo-hi-lo lo-lo-hi-lo
	430433	AA478883	Hs.273766	ESTs	lo-lo-hi-lo
	435351	T80177	Hs.118064	similar to rat nuclear ubiquitous casein	lo-lo-hi-lo
	403218	AL134878		ribosomal protein, large P2	lo-lo-hi-lo
~~	420678	AW593288	Hs.3530	TLS-associated serine-arginine protein 2	lo-lo-hi-lo
75	445808	AV655234		ESTs, Moderately similar to PC4259 ferri	lo-lo-hi-lo
	429933	AA765596	Hs.187691	ESTs	lo-lo-hi-lo
	419802	AA250950	Hs.154334	ESTs	lo-lo-hi-lo
	425155	W26522	Hs.75890	gb:32g2 Human retina cDNA randomly prime	lo-lo-hi-lo
80	417314	N68168 Al932995	Hs.183475	gb:za11c01.s1 Soares fetal liver spleen Homo sapiens clone 25061 mRNA sequence	lo-lo-hi-lo
30	428290 422128	AW881145	16.105473	gb:QV0-OT0033-010400-182-207 OT0033 Homo	lo-lo-hi-lo lo-lo-hi-lo
	432014	H66741	Hs.38540	ESTs, Weakly similar to ALU4_HUMAN ALU S	lo-lo-hi-lo lo-lo-hi-lo
	.02917			==, w/mor_stodest ALO d	טרוורטיייי

	407351	AW383165		gb:PM3-HT0344-151299-004-f07 HT0344 Hamo	lo-lo-hi-lo
	443231	W87548	Hs.132932	ESTs	lo-lo-hi-lo
	444001	AI095087	Hs.152299	ESTs, Moderately similar to S65657 alpha	lo-lo-hi-lo
_	435064		Hs.31433	ESTs	lo-lo-hi-lo
5	435173		Hs.255451	ESTs	lo-lo-hi-lo
	411831	AW994394		gb:RC3-BN0036-060400-014-h12 BN0036 Homo	lo-lo-hi-lo
	446572	AV659151	Hs.282961	ESTs	lo-lo-hi-lo
	428114	A1821548	Hs.98363	ESTs, Wealdy similar to I38022 hypotheti	lo-lo-hi-lo
10	406207			Target Exon	lo-lo-hi-lo
10	405011			Target Exon	lo-lo-hi-lo
	409451	AF012626	Hs.54472	fragile X mental retardation 2	lo-lo-hi-lo
	411233	AW833793		gb:QV4-TT0008-130100-080-a06 TT0008 Homo	lo-lo-hi-lo
	455729	BE072092		gb:PM4-BT0532-160200-003-b11 BT0532 Homo	lo-lo-hi-lo
	439454	AA836120	Hs.258958	ESTs	lo-lo-hi-lo
15	445124	A1806403	Hs.143942	ESTs	lo-lo-hi-lo
	410324	AW292539	Hs.30177	ESTs	lo-lo-hi-lo
	446548	AI769392	Hs.200215	ESTs	lo-lo-hi-lo
	416999	AW195747	Hs.21122	hypothetical protein FLJ11830 similar to	lo-lo-hi-lo
	414553	Al813865	Hs.164478	hypothetical protein FLJ21939 similar to	lo-lo-hi-lo
20	444647	H14718	Hs.11506	Human clone 23589 mRNA sequence	lo-lo-hi-lo
	418271	NM_000919		peptidylglycine alpha-amidating monooxyg	lo-lo-hi-lo
	407939	W05608	Hs.312679	ESTs, Weakly similar to A49019 dynein he	lo-lo-hi-lo
	432676	AJ187366	113.012013	gb:qf29c01.x1 Soares_testis_NHT Homo sap	
			U= 70000		lo-lo-hi-lo
25	415156	X84908	Hs.78060	phosphorylase kinase, beta	lo-lo-hi-lo
25	432679	Al146956	Hs.146723	ESTs, Weakly similar to A53950 transcrip	lo-lo-hi-lo
	412121	AB033061	Hs.73287	KIAA1235 protein	lo-lo-hi-lo
	418858	AW961605	Hs.21145	hypothetical protein RG083M05.2	lo-lo-hi-lo
	425204	NM_002436		membrane protein, palmitoylated 1 (55kD)	lo-lo-ni-lo
20	418348	A1537167	Hs.96322	hypothetical protein FLJ23560	lo-lo-hi-lo
30	410765	A1694972	Hs.66180	nucleosome assembly protein 1-like 2	lo-lo-hi-lo
	445594	AW058463	Hs.12940	zinc-fingers and homeoboxes 1	lo-lo-hi-lo
	416503	H98502	Hs.269853	ESTs	lo-lo-hi-lo
	426167	AF039023	Hs.167496	RAN binding protein 6	lo-lo-hi-lo
	451752	AB032997.	Hs.26966	KIAA1171 protein	lo-lo-hi-lo
35	447124	AW976438	Hs.17428	RBP1-like protein	lo-lo-hi-lo
	419872	Al422951	Hs.146162	ESTs	lo-lo-hi-lo
	443161	Al038316		gb:ox48c08.x1 Soares_total_fetus_Nb2HF8_	lo-lo-hi-lo
	445391	T92576	Hs.191168	ESTs	lo-lo-hi-lo
	443801	AW206942	Hs.253594	intron of: trichorhinophalangeal syndro	lo-lo-hi-lo
40	446706	AW807631	Hs.190488	Homo sapiens, Similar to nuclear localiz	lo-lo-hì-lo
. •	428172	U09367	Hs.182828	zinc finger protein 136 (clone pHZ-20)	lo-lo-hi-lo
	421021	AA808018	Hs.109302	ESTs	lo-lo-hi-lo
	431749	AL049263	Hs.306292	Homo sapiens mRNA; cDNA DKFZp564F133 (fr	lo-lo-hi-lo
	423784	AK000039	Hs.132826	Homo sapiens cDNA FLJ14913 fis, clone PL	lo-lo-hi-lo
45	419479	A1288348	Hs.23450	milochondrial ribosomal protein S25	lo-lo-hi-lo
73	450900	H61005	Hs.37902	ESTs	lo-lo-hi-lo
	423396	AI382555	Hs.127950	bromodomain-containing 1	lo-lo-hi-lo
	426137	AL040683	Hs.167031	DKFZP566D133 protein	lo-lo-hi-lo
50	442012	A1733277	Hs.128321	ESTs	lo-lo-hi-lo
50	452271	AA025976	Hs.34569	ESTs	lo-lo-hi-lo
	414882	D79994	Hs.77546	Homo sapiens cDNA: FLJ21983 fis, clone H	ło-lo-hi-lo
	432195	AJ243669	Hs.8127	KIAA0144 gene product	lo-lo-hi-lo
	430217	N47863	Hs.180450	ribosomal protein S24	lo-lo-hi-lo
55	429567	R35606	Hs.326800	Human EST clone 53125 mariner transposon	lo-lo-hi-lo
22	438810	AW897846	Hs.6421	hypothetical protein DKFZp761N09121	lo-lo-hi-lo
	436796	BE515260	Hs.5320	hypothetical protein	lo-lo-hi-lo
	426352	N72324	Hs.55098	ESTs	io-io-hi-lo
	415308			gb:HSC04H101 normalized Infant brain cDN	lo-lo-hi-lo
60	420148	U34227	Hs.95361	myosin VIIA (Usher syndrome 1B (autosoma	io-lo-hi-lo
60	434442	AA737415	Hs.152826	ESTs	lo-lo-hi-lo
	449429	AA054224	Hs.59847	ESTs	lo-lo-hi-lo
	410245	C17908	Hs.194125	ESTs	lo-lo-hi-lo
	421168	AF182277	Hs.330780	cytochrome P450, subfamily IIB (phenobar	lo-lo-hi-lo
	436237	R11528	Hs.271968	ESTs	io-lo-hi-lo
65	440668	A1989538	Hs.191074	ESTs	lo-lo-hi-lo
	422068	AI807519	Hs.104520	Homo sapiens cDNA FLJ13694 fis, clone PL	lo-lo-hi-lo
	410216	BE061839		gb:RC1-BT0254-290100-015-a05 BT0254 Homo	lo-lo-hi-lo
	439437	A1207788	Hs.343628	sialyltransferase 4B (beta-galactosidase	lo-lo-hi-lo
	417061	AI675944	Hs.188691	Homo sapiens cDNA FLJ12033 fis, clone HE	to-lo-hi-lo
70	403046			NM_005656*:Homo sapiens transmembrane pr	lo-lo-hi-lo
	404528	AI912555		peptide YY, 2 (seminatplasmin)	lo-lo-hi-lo
	439734	AC005013	Hs.149	cAMP response element-binding protein CR	lo-lo-hi-lo
	452997	N64777	Hs.44656	ESTs	ko-lo-hi-lo
	403745		1 ,000	ENSP00000226812*:KIAA1494 protein (Fragm	lo-lo-hi-lo
75	411448	AA178955	Hs.271439	ESTs, Weakly similar to 138022 hypotheti	io-lo-hi-lo
, ,	422460	AW445014	Hs.197746	ESTs Veakly sirillar to 130022 hypotheti	lo-lo-hi-lo
		M11440014	110.13/140		
	404058	DE15/087	He 136660	Target Exon ESTs. Weathy similar to 7N91. HI IMAN 7INC	lo-lo-hi-lo
	436184	BE154067	Hs.136660	ESTs, Weakly similar to ZN91_HUMAN ZINC	lo-lo-hi-lo
80	427702	N76589	Hs.14454	ESTs, Wealdy similar to TFIID subunit TA	lo-lo-hi-lo
OU	440695	AW088363	Hs.246240	ESTS	lo-lo-hi-lo
	424881	AL119690	Hs.153618	HCGVIII-1 protein	lo-lo-hi-hi
	440573	BE550891	Hs.270624	ESTs	lo-lo-hi-hi
				150	

	*****	1100040			
	416659	W22048	Hs.64753	gb:61A12 Human relina cDNA Tsp509I-cleav	lo-lo-hì-hi
	436731	AA580691	Hs.180789	S164 protein	lo-lo-hi-hi
	405102	*1000000	11 004004	C15001220*:gi]4469558[gb]AAD21311.1] (AF	lo-lo-hi-hi
5	450219	A1826999	Hs.224624	ESTs	lo-lo-hi-hi
J	404527	Al912555	11 400000	peptide YY, 2 (seminalplasmin)	lo-lo-hi-hi
	439158	R60323	Hs.193888	ESTs	lo-lo-hi-hi
	431952	Z70695	Hs.272240	Homo sapiens cDNA FLJ11086 fis, clone PL	lo-lo-hi-hi
	418584	NM_004606	Hs.182339	TATA box binding protein (TBP)-associate	lo-lo-hi-hi
10	424241 410124	AW995948 AW962229	Hs.128927	Homo sapiens pyruvate dehydrogenase kina Homo sapiens cDNA FLJ13903 fis, clone TH	lo-lo-hi-hi lo-lo-hi-hi
10	435955	AA830515	Hs.222917	ESTs	lo-lo-hi-hi
	424001	W67883	Hs.137476	paternally expressed 10	hi-hi-to-to
	441399	A1630844	Hs.126919	ESTs	hi-hi-to-to
	440184	AB002297	Hs.7022	dedicator of cyto-kinesis 3	hi-hi-lo-lo
15	421996	AW583807	Hs.1460	glucagon	hi-hi-lo-lo
10	444252	R21135	Hs.54985	ESTs `	hi-hi-lo-lo
	402082	141100	1.5.5 .555	C18000743*:gi]6678363[ref]NP_033416.1] t	hi-hi-lo-lo
	405396			C22000452*:gi[6981522]ref[NP_036781.1] r	hi-hi-lo-lo
	412457	T32587	Hs.170414	paired basic amino acid cleaving system	hi-hi-lo-lo
20	415808	R21439	Hs.334578	Homo sapiens, clone IMAGE:3929520, mRNA	hi-hi-lo-lo
	441494	AW452344	Hs.129977	ESTs	hi-hi-lo-lo
	437330	AL353944	Hs.50115	Homo sapiens mRNA; cDNA DKFZp761J1112 (f	hi-hi-lo-lo
	452784	BE463857	Hs.151258	hypothetical protein FLJ21062	hi-hi-lo-lo
	410037	AB020725	Hs.58009	KIAA0918 protein	hi-hi-lo-lo
25	449145	Al632122	Hs.198408	ESTs	hi-hi-lo-lo
	452487	AW207659	Hs.6630	Homo sapiens cDNA FLJ13329 fis, clone OV	hi-hi-lo-lo
	431031	AA830335	Hs.105273	ESTs	hi-hi-lo-lo
	427209	H06509	Hs.92423	KIAA1566 prolein	hi-hi-lo-lo
	434280	BE005398		gb:CM1-BN0116-150400-189-h02 BN0116 Homo	hi-hi-lo-lo
30	418236	AW994005	Hs.337534	ESTs	hi-hi-lo-lo
	429201	X03178	Hs.198246	group-specific component (vitamin D bind	hi-hi-lo-lo
	416653	AA768553	Hs.193145	metallothionein 1E (functional)	hi-hi-lo-lo
	422501	AA354690	Hs.144967	ESTs	hi-hi-lo-lo
25	425087	R62424	Hs.126059	ESTs	hi-hi-lo-lo
35	426798	AA385062	Hs.130260	ESTs	hi-hi-lo-lo
	443798	R07848	Hs.188522	ESTs	hi-hi-lo-lo
	427254	AL121523	Hs.97774	ESTs	hi-hi-lo-lo
	431657	Al345227	Hs.105448	ESTs, Weakly similar to B34087 hypotheti	hi-hi-lo-lo
40	409963	AA133590	Hs.250857	calcium/calmodulin-dependent protein kin	hi-hi-lo-lo
40	446006	NM_004403	HS.13530	deafness, autosomal dominant 5	hi-hi-lo-lo
	418259	AA215404	Un 440044	ESTs	hi-hi-lo-lo
	410173	AA706017	Hs.119944	ESTs	hi-hi-lo-lo
	436023 448428	T81819 AF282874	Hs.302251 Hs.21201	ESTs	hi-hi-lo-lo
45	430665	BE350122	Hs.157367	nectin 3; DKFZP566B0846 protein ESTs, Wealdy similar to 178885 serine/th	hi-hi-lo-lo hi-hi-lo-lo
73	432559	AW452948	Hs.257631	ESTs	hi-hi-lo-lo
	451572	AA018556	Hs.268691	ESTs, Moderately similar to ALU2_HUMAN A	hi-hi-lo-lo
	456032	AW957446	Hs.301711	ESTs	hi-hi-lo-lo
	438209	AL120659	Hs.6111	aryl-hydrocarbon receptor nuclear transl	hi-hi-lo-lo
50	438337	AK002058	Hs.6166	hypothetical protein FLJ11196	hi-hi-lo-lo
	431795	AK002088	Hs.270124	Homo sapiens cDNA FLJ11226 fis, clone PL	hi-hi-lo-lo
	421114	AW975051	Hs.293156	ESTs, Weakly similar to 178885 serine/th	hi-hi-lo-lo
	431843	AA516420		ESTs, Weakly similar to 138022 hypotheti	hi-hi-lo-lo
	440948	AW188311	Hs.128619	ESTs	hi-hi-lo-lo
55	430105	X70297	Hs.2540	cholinergic receptor, nicotinic, alpha p	hi-hi-lo-lo
	439046	AA947354		gb:od86e11.s1 NCI_CGAP_Ov2 Homo sapiens	hi-hi-lo-lo
	451491	A1972094	Hs.286221	Homo sapiens cDNA FLJ13741 fis, clone PL	hi-hi-lo-lo
	452789	AW081626	Hs.242561	ESTs	hi-hi-lo-lo
60	419829	A1924228	Hs.115185	ESTs, Moderately similar to PC4259 ferri	hi-hi-lo-lo
60	449567	A1990790	Hs.188614	ESTs	hi-hi-lo-lo
	407787	N21307	Hs.13477	ESTs, Weakly similar to 1207289A reverse	hi-hi-lo-lo
	409091	AW970386	Hs.269423	ESTs	hi-hi-lo-lo
	435354	AA678267	Hs.117115	ESTs	hi-hi-lo-lo
65	444809	BE207568	Hs.208219	oculospanin	hi-hi-lo-lo
05	422170	AI791949	Hs.112432	anti-Mullerian hormone	hi-hi-lo-lo
	453582	AW854339	Hs.33476	hypothetical protein FLJ11937	hi-hi-lo-lo
	435905	AW997484	Hs.5003	KIAA0456 protein	hi-hi-lo-lo
	443884	N20617 AB023197	Hs.194397	leptin receptor	hi-hi-lo-lo hi-hi-lo-lo
70	430027 432582	AJ623817	Hs.227743 Hs.168457	KIAA0980 protein ESTs	hi-hi-lo-to
, 0	417993	AW963705	Hs.301183	molecule possessing ankyrin repeals indu	hi-hi-lo-lo
	444930	BE185536	Hs.301183	molecule possessing ankyrin repeats indu	hi-hi-lo-lo
	427794	AA709186	Hs.99070	ESTs	hi-hi-lo-lo
	410913	AL050367	Hs.66762	Homo sapiens mRNA; cDNA DKFZpS64A026 (fr	hi-hi-lo-lo
75	431992	NM_002742		prolein kinase C, mu	hi-hi-lo-lo
	447846	AA324057	Hs.77955	Homo sapiens cDNA: FLJ23527 fis, clone L	hi-hi-lo-lo
	430439	AL133561		DKFZP434B061 protein	hi-hi-lo-lo
	432621	Al298501	Hs.12807	ESTs, Weakly similar to T46428 hypotheti	hi-hi-lo-lo
0.0	431427	AK000401	Hs.252748	Homo sapiens cDNA FLJ20394 fis, clone KA	hi-hi-lo-lo
80	408872	Al476139	Hs.13291	ESTs	hi-hi-lo-lo
	453200	AA033832	Hs.212433	ESTs	hi-hi-lo-lo
	411529	AA430348	Hs.317596	Homo sapiens cDNA FLJ12927 fis, clone NT	hi-hi-lo-lo
				4.004	

	414483	R25513	Hs.10683	ESTs	hi-hi-lo-lo
	451273	NM_014811		KIAA0649 gene product	hi-hi-lo-lo
	437052	AA861697	Hs.120591	ESTs	hi-hi-lo-lo
5	440049	R06699	Hs.19769	hypothetical protein MGC4174	hi-hi-lo-lo
5	429483	AA974832	Hs.128708	ESTs	hi-hi-lo-lo
	411296 425188	BE207307 AK002052	Hs.10114 Hs.155071	growth suppressor 1 hypothetical protein FLJ11190	hi-hi-lo-lo hi-hi-lo-lo
	436315	BE390513	Hs.27935	hypothetical protein MGC4837	hi-hi-lo-lo
	400297	Al127076	Hs.306201	hypothetical protein DKFZp564O1278	hi-hi-lo-lo
10	431089	BE041395	113.000201	ESTs, Weakly similar to unknown protein	hi-hi-lo-lo
10	418824	AW751661	Hs.53542	choreoacanthocytosis gene; KIAA0986 prot	hi-hi-lo-lo
	449226	AB002365	Hs.23311	KIAA0367 protein	hi-hi-lo-lo
	450149	AW969781	Hs.132863	Zic family member 2 (odd-paired Drosophi	hi-hi-lo-lo
	418443	NM_005239	Hs.85146	v-ets avian erythroblastosis virus E26 o	hi-hi-lo-lo
15	458692	BE549905	Hs.231754	ESTs	hi-hi-lo-lo
	410102	AW248508	Hs.279727	ESTs; homologue of PEM-3 [Ciona savignyi	hi-hi-lo-lo
	451062	AL110125	Hs.25910	Homo sapiens mRNA; cDNA DKFZp564C1416 (f	hi-hi-lo-lo
	407633	NM_007069		similar to rat HREV107	hi-hi-lo-lo
20	418941	AA452970	Hs.239527	E1B-55kDa-associated protein 5	hi-hi-lo-lo
20	407059	X95406		gb:H.sapiens cyclin E gene.	hi-hi-lo-lo
	455956	BE162704	11- 5024	gb:PM1-HT0454-301299-001-d08 HT0454 Homo	hi-hi-lo-lo
	437763	AA469369	Hs.5831	tissue inhibitor of metalloproteinase 1	hi-hi-lo-lo hi-hi-lo-lo
	451404	AA460775	Hs.6295	ESTs, Wealdy similar to T17248 hypotheti	
25	428494 414957	AA233439 D61283	Hs.184634 Hs.45206	hypothetical protein FLJ20005 ESTs	hi-hi-lo-lo hi-hi-lo-lo
23	456415	Al734051	Hs.277102	ESTs, Weakly similar to ALU1_HUMAN ALU S	hi-hi-lo-lo
	400183	A1734031	113.211102	Eos Control	hi-hi-lo-lo
	400158			ENSP00000244302*:CDNA FLJ11591 fis, clon	hi-hi-lo-lo
	403893			ENSP00000237068*:Prolocadherin alpha 6 p	hi-hi-lo-lo
30	423809	AJ223833	Hs.154483	ESTs	hi-hi-lo-lo
	400170			Eos Control	hi-hi-lo-lo
	403291			Target Exon	hi-hi-lo-lo
	422026	UB0736	Hs.110826	trinucleotide repeat containing 9	hi-hi-lo-lo
~~	417130	AW276858	Hs.81256	S100 calcium-binding protein A4 (calcium	hi-hi-lo-lo
35	432472	AA548781	Hs.136418	ESTs	hi-hi-lo-lo
	405231			C2001066:gi 10257425 ref NP_033892.1  CD	hi-hi-lo-lo
	400141			Eos Control	hi-hi-lo-lo
	428971	BE278404	Hs.285813	hypothetical protein FLJ11807	hi-hi-lo-lo
40	422390	AW450893	Hs.121830	ESTs, Weakly similar to T42682 hypotheti	hi-hi-lo-lo
40	425538	BE270918	Hs.164026	Horno sapiens, clone IMAGE:3534875, mRNA,	hi-hi-lo-lo
	456972	AI054347	Hs.2017	ribosomal protein L38	hi-hi-lo-lo
	456622	AF205849	Hs.107740	Kruppel-like factor 2 (lung)	hi-hi-lo-lo
	418515 448439	Al568453 BE613082	Hs.19487 Hs.28229	ESTs, Weakly similar to CNIH_HUMAN CORNI	hi-hi-lo-lo hi-hi-lo-lo
45	445418	AW139377	Hs.127179	ARG99 protein cryptic gene	hi-hi-lo-lo
73	402559	Z23024	115.121113	Rho GTPase activating protein 1	hi-hi-lo-lo
	402575	223024		Rho GTPase activating protein 1	hi-hi-lo-lo
	420811	AA807544		ESTs, Weakly similar to 834323 GTP-bindi	hi-hi-lo-lo
	446627	Al973016	Hs.15725	hypothetical protein SBBI48	hi-hi-lo-lo
50	400247			Eos Control	hi-hi-lo-lo
	430289	AK001952	Hs.238039	hypothetical protein FLJ11090	hi-hi-lo-lo
	400133			Eos Control	hi-hi-lo-lo
	418816	T29621	Hs.88778	carbonyl reductase 1	hi-hi-lo-lo
<i>5 5</i>	433579	BE264473	Hs.284297	hypothetical protein from EUROIMAGE 1967	hi-hi-lo-lo
55	401952	***********		Target Exon	hi-hi-lo-lo
	410349	AW663021	Hs.323445	ESTs, Weakly similar to T2D3_HUMAN TRANS	hi-hi-lo-lo
	417558	AF045229	Hs.82280	regulator of G-protein signalling 10	hi-hi-lo-lo
	446851 404489	AW007332	Hs.10450	Homo sapiens cDNA: FLJ22063 fis, clone H Target Exon	hi-hi-lo-lo hi-hi-lo-lo
60	405802			Target Exon	hi-hi-lo-lo
00	456266	L29073	Hs.198726	cold shock domain protein A	hi-hi-lo-lo
	457133	M54968		v-Ki-ras2 Kirsten rat sarcoma 2 viral on	hi-hi-lo-lo
	459330	C16931		gb:C16931 Clontech human aorta polyA mRN	hi-hi-lo-lo
	433041	BE265848	Hs.289080	colon cancer-associated protein Mic1	lo-to-lo-hi
65	446545	Al431798	Hs.164192	ESTs, Weakly similar to Y161_HUMAN HYPOT	lo-lo-lo-hi
	414911	NM_000107	Hs.77602	damage-specific DNA binding protein 2 (4	lo-lo-lo-hi
	414682	AL021154	Hs.76884	inhibitor of DNA binding 3, dominant neg	lo-lo-lo-hi
	422311	AF073515	Hs.114948	cytokine receptor-like factor 1	lo-lo-lo-hi
70	447329	BE090517		ESTs, Moderately similar to ALU8_HUMAN A	lo-lo-lo-hi
70	412942	AL120344	Hs.75074	milogen-activated protein kinase-activat	lo-lo-lo-hi
	420747	BE294407	Hs.99910	phosphofructokinase, platelet	lo-lo-lo-hi
	431912	AI660552	Hs.76549	ESTs, Weakly similar to A56154 Abl subst	lo-lo-lo-hi
	446506	Al123118	Hs.15159	chemokine-like factor, alternatively spl	lo-lo-lo-hi lo-lo-lo-hi
75	408633	AW963372	Hs.46677	PRO2000 protein	10-10-10-11 hi-lo-lo-hi
13	433675	AW977653	Hs.75319 Hs.150555	ribonucleotide reductase M2 polypeptide protein predicted by clone 23733	nı-to-to-nı hi-lo-lo-hi
	424560 425234	AA158727 AW152225	Hs.165909	ESTs, Weakly similar to 138022 hypotheti	ni-to-to-ni hi-to-to-hi
	425254	AA206079	Hs.6693	hypothetical protein FLJ20420	hi-lo-lo-hi
	410174	AA306007	Hs.59461	DKFZP434C245 protein	hi-to-lo-hi
80	410442	X73424	Hs.63788	propionyl Coenzyme A carboxylase, beta p	hi-lo-lo-hi
	429190	H18650	Hs.92602	ESTs	hi-lo-lo-hi
	423619	T48691	Hs.249159	adrenergic, alpha-2A-, receptor	hi-lo-lo-hi
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	400304	********	11- 20000	rot-	
	433764	AW753676	Hs.39982	ESTS	hi-lo-lo-hi
	421998	R74441	Hs.117176	poly(A)-binding protein, nuclear 1	hi-lo-lo-hi
	451593	AF151879	Hs.26706	CGI-121 protein	hi-lo-lo-hi
5	452092	BE245374	Hs.27842	hypothetical protein FLJ11210	hi-lo-lo-hi
5	447425	AI963747	Hs.18573	acylphosphatase 1, erythrocyte (common)	hi-lo-lo-hi
	421654	AW163267	Hs.106469	suppressor of var1 (S.cerevisiae) 3-like	hi-lo-lo-hi
	432502	NM_014641		KIAA0170 gene product	hi-lo-lo-hi
	429597	NM_003816		a disintegrin and metalloproteinase doma	hi-lo-lo-hi
10	434203	BE262677	Hs.283558	hypothetical protein PRO1855	hi-lo-lo-hi
10	438461	AW075485	Hs.286049	phosphoserine aminotransferase	hi-lo-lo-hi
	409142	AL136877	Hs.50758	SMC4 (structural maintenance of chromoso	hi-lo-lo-hi
	439574 438182	AJ469788	Hs.165190 Hs.182545	ESTS Whooldy cimiles to Al 114 Little AN ALLI C	hi-lo-lo-hi
	449103	AW342140 T24968	Hs.23038	ESTs, Weakly similar to ALU1_HUMAN ALU S	hi-lo-lo-hi
15		Al654133		HSPC071 prolein	hi-lo-lo-hi
13	421059		Hs.30212	thyroid receptor interacting protein 15	hi-lo-lo-hi
	446939	AL133353	Hs.16606	CGI-32 protein	hi-lo-lo-hi
	408576 410073	NM_003542		H4 histone family, member G	hi-lo-lo-hi
		AW408163	Hs.58488	catenin (cadherin-associated protein), a	hi-lo-lo-hi
20	450912	AW939251	Hs.25647	v-fos FBJ murine osteosarcoma viral onco	hi-lo-lo-hi
20	434701 450455	AA460479	Hs.321707	KIAA0742 protein	hi-lo-lo-hi
	451144	AL117424 AW956103	Hs.25035 Hs.61712	chloride intracellular channel 4	hi-lo-lo-hi
	427390	AW930103 AI432163	Hs.268231	pyruvate dehydrogenase kinase, isoenzyme	hi-lo-lo-hi
	451831	NM_001674		Homo sapiens cDNA: FLJ23111 fis, clone L	hi-lo-lo-hi
25	406776	T16206	Hs.237164	activating transcription factor 3	hi-lo-lo-hi
23	428157	Al738719	Hs.198427	ESTs, Highly similar to LDHH_HUMAN L-LAC hexokinase 2	hi-lo-lo-hi
	408096	BE250162	Hs.83765	dihydrofolate reductase	hi-lo-lo-hi
	418203	X54942	Hs.83758	CDC28 protein kinase 2	hi-lo-lo-hi
	449338	H73444	Hs.394	adrenomedullin	hi-lo-lo-hi
30	422082	AA016188	Hs.111244	hypothetical protein	hi-lo-lo-hi
50	407907	AI752235	Hs.41270	procollagen-lysine, 2-oxoglutarate 5-dio	hi-lo-lo-hi hi-lo-lo-hi
	416655	AW968613	Hs.79428	BCL2/adenovirus E1B 19kD-interacting pro	
	419551	AW582256	Hs.91011	anterior gradient 2 (Xenepus laevis) hom	hi-lo-lo-hi hi-lo-lo-hi
	434094	AA305599	Hs.238205	hypothetical protein PRO2013	hi-lo-lo-hi
35	443951	F13272	Hs.111334	femilin, light polypeptide	
55	422975	AA347720	Hs.122669	KIAA0264 protein	hi-lo-lo-hi hi-lo-lo-hi
	430314	AA369601	Hs.239138	pre-B-cell colony-enhancing factor	hi-lo-lo-hi
	412664	AA421404	Hs.346868	nucleolar protein p40; homolog of yeast	hi-lo-lo-hi
	408089	H59799	Hs.42644	thioredoxin-like	hi-lo-lo-hi
40	409690	W45393	Hs.55888	activating transcription factor 7	hi-lo-lo-hi
••	442332	Al693251	Hs.8248	Target CAT	hi-lo-lo-hi
	408388	AF091086	Hs.44563	hypothetical protein	hi-lo-lo-hi
	441252	AW360901	Hs.183047	hypothetical protein MGC4399	hi-lo-lo-hi
	433069	X76732	Hs.3164	nucleobindin 2	hi-lo-lo-hi
45	443837	AI984625	Hs.9884	spindle pole body protein	hi-lo-lo-hi
	426108	AA622037	Hs.166468	programmed cell death 5	hi-lo-lo-hi
	441181	AA416925	Hs.121076	peptidylprolyl isomerase (cyclophilin)-l	hi-lo-lo-hi
	447397	BE247676	Hs.18442	E-1 enzyme	hi-lo-lo-hi
	427505	AA361562	Hs.178761	26S proleasome-associated pad1 homolog	hi-lo-lo-hi
50	430287	AW182459	Hs.125759	ESTs, Weakly similar to LEU5_HUMAN LEUKE	hi-lo-lo-hi
	415857	AA866115	Hs.127797	Homo sapiens cDNA FLJ11381 fis, clone HE	hi-lo-lo-hi
	423198	M81933	Hs.1634	cell division cycle 25A	hi-lo-lo-hi
	407687	AK002011	Hs.37558	hypothetical protein FLJ11149	hi-lo-lo-hi
	431374	BE258532	Hs.251871	CTP synthase	hi-lo-lo-hi
55	413273	U75679	Hs.75257	stem-loop (histone) binding protein	hi-lo-lo-hi
	442799	Al564739	Hs.68505	ESTs	hi-lo-lo-hi
	443881	R64512	Hs.237146	hypothetical protein FLJ12752	hi-lo-lo-hi
	416209	AA236776	Hs.79078	MAD2 (mitotic arrest deficient, yeast, h	hi-lo-lo-hi
<i>(</i> 0	421834	BE543205	Hs.288771	DKFZP586A0522 protein	hi-lo-lo-hi
60	411263	BE297802	Hs.69360	kinesin-like 6 (mitotic centromere-assoc	hi-lo-lo-hi
	413924	AL119964	Hs.75616	seladin-1	hi-lo-lo-hi
	450598	AF151076	Hs.25199	hypothetical protein	hi-lo-lo-hi
	439453	BE264974	Hs.6566	thyroid hormone receptor interactor 13	hi-lo-lo-hi
CE	429612	AF062649	Hs.252587	pituitary tumor-transforming 1	hi-lo-lo-hi
65	443426	AF098158	Hs.9329	chromosome 20 open reading frame 1	hi-lo-lo-hi
	452353	C18825	Hs.29191	epithelial membrane protein 2	hi-lo-lo-hi
	419879	Z17805	Hs.93564	Homer, neuronal immediate early gene, 2	hi-lo-lo-hi
	422363	T55979	Hs.115474	replication factor C (activator 1) 3 (38	hi-lo-lo-hi
70	416065	BE267931	Hs.78996	proliferating cell nuclear antigen	hi-lo-lo-hi
70	424308	AW975531	Hs.154443	minichromosome maintenance deficient (S.	hi-lo-lo-hi
	447519	U46258	Hs.339665	ESTs	hi-lo-lo-hi
	437679	NM_014214		inositol(myo)-1(or 4)-monophosphatase 2	hi-lo-lo-hi
	446636	AC002563	Hs.15767	citron (rho-interacting, serine/threonin	hi-lo-lo-hi
75	422094	AF129535	Hs.272027	F-box only protein 5	hi-lo-lo-hi
75	440334	BE276112	Hs.7165	zinc finger protein 259	hi-lo-lo-hi
	421921	H83363	Hs.6820	translocase of inner mitochondrial membr	hi-lo-lo-hi
	422938	NM_001809		centromere protein A (17kD)	hi-lo-lo-hi
	427719	AJ393122	Hs.134726	ESTs	hi-lo-lo-hi
80	422283	AW411307	Hs.114311	CDC45 (cell division cycle 45, S.cerevis	hi-lo-lo-hi
OU	424840	D79987	Hs.153479	extra spindle poles, S. cerevisiae, homo	hi-lo-lo-hi
	418216	AA662240	Hs.283099	AF15q14 protein	hi-lo-lo-hi
	412140	AA219691	Hs.73625	RAB6 interacting, kinesin-like (rabkines	hi-lo-lo-hi
				172	

	418322	AA284166	Hs.84113	cyclin-dependent kinase inhibitor 3 (CDK	hi-lo-lo-hi
	428479	Y00272	Hs.334562	cell division cycle 2, G1 to S and G2 to	hi-lo-lo-hi
	449722	BE280074	Hs.23960	cyclin B1	hi-lo-lo-hi
5	417933 433001	X02308	Hs.82962	thymidylate synthetase	hi-lo-lo-hi
,	413943	AF217513 AW294416	Hs.279905 Hs.144687	clone HQ0310 PRO0310p1 Homo sapiens cDNA FLJ12981 fis, clone NT	hi-lo-lo-hi hi-lo-lo-hi
	424905		Hs.153704	NIMA (never in mitosis gene a)-related k	hi-lo-lo-hi
	422765	AW409701	Hs.1578	baculoviral IAP repeat-containing 5 (sur	hi-lo-lo-hi
	425397	J04088	Hs.156346	topoisomerase (DNA) II alpha (170kD)	hi-lo-lo-hi
10	444371	BE540274	Hs.239	forkhead box M1	hi-lo-lo-hi
	422956	BE545072	Hs.122579	ECT2 protein (Epithelial cell transformi	hì-lo-lo-hi
	444783	AK001468	Hs.62180	anillin (Drosophila Scraps homolog), act	hi-lo-lo-hi
	453884	AA355925	Hs.36232	KIAA0186 gene product	hi-lo-lo-hi
1.5	416980	AA381133	Hs.80684	high-mobility group (nonhistone chromoso	hi-lo-lo-hi
15	442432	BE093589	Hs.38178	hypothetical protein FLJ23468	hi-lo-lo-hi
	417308	H60720	Hs.81892	KIAA0101 gene product	hi-lo-lo-hi
	433133	AB027249	Hs.104741	PDZ-binding kinase; T-cell originated pr	hi-lo-lo-hi
	432626	AA471098	Hs.278544	acetyl-Coenzyme A acetyltransferase 2 (a	hi-lo-lo-hi
20	441020 412281	W79283 Al810054	Hs.35962 Hs.14119	ESTs ESTs	hi-lo-lo-hi
20	435602	AF217515	Hs.283532	uncharacterized bone marrow protein BM03	hi-lo-lo-hi hi-lo-lo-hi
	400882	W 511210	115.205552	Target Exon	hi-lo-lo-hi
	446269	AW263155	Hs.14559	hypothetical protein FLJ10540	hi-lo-lo-hi
	417847	Al521558	Hs.7331	hypothetical protein FLJ22316	hi-lo-lo-hi
25	400881	, 402 1000	. 10.7 00 1	NM_025080:Homo sapiens hypothetical prot	hi-lo-lo-hi
	419356	AI656166	Hs.7331	hypothetical protein FLJ22316	hi-lo-lo-hi
	400292	AA250737	Hs.72472	BMP-R1B	hi-lo-lo-hi
	415539	AI733881	Hs.72472	BMP-R1B	hi-lo-lo-hi
20	453935	Al633770	Hs.42572	ESTs	hi-lo-lo-hi
30	420005	AW271106	Hs.133294	ESTs	hi-lo-lo-hi
	428450	NM_014791		KIAA0175 gene product	hi-lo-lo-hi
	436291	BE568452	Hs.344037	protein regulator of cytokinesis 1	hi-lo-lo-hi
	441362	BE614410	Hs.23044	RAD51 (S. cerevisiae) homolog (E coli Re	hi-lo-lo-hi
35	428484 418526	AF104032 BE019020	Hs.184601 Hs.85838	solute carrier family 7 (cationic amino .	hì-lo-lo-hi
55	458809	AW972512	Hs.20985	solute carrier family 16 (monocarboxylic sin3-associated polypeptide, 30kD	hi-lo-lo-hi hi-lo-lo-hi
	444984	H15474	Hs.132898	fally acid desalurase 1	hi-lo-lo-hi
	447342	Al199268	Hs.19322	Homo sapiens, Similar to RIKEN cDNA 2010	hi-hi-lo-lo
	428330	L22524	Hs.2256	matrix metalloproteinase 7 (matrilysin,	hi-hi-lo-lo
40	428336	AA503115	Hs.183752	microserninoprotein, beta-	hi-hi-lo-lo
	430389	AL117429	Hs.240845	DKFZP434D146 protein	hi-hi-lo-lo
	417318	AW953937	Hs.240845	ESTs	hi-hi-lo-lo
	422545	X02761	Hs.287820	fibronectin 1	hi-hi-lo-lo
45	417640	D30857	Hs.82353	protein C receptor, endothelial (EPCR)	hi-lo-lo-lo
45	422809	AK001379	Hs.121028	hypothetical protein FLJ10549	hi-lo-lo-hi
	425580	L11144	Hs.1907	galanin	hi-lo-io-hi
	416836	D54745	Hs.80247	cholecystokinin	hi-lo-lo-hi
	434170 427958	AA626509 AA418000	Hs.122329 Hs.98280	ESTs potassium intermediate/small conductance	hi-io-io-hi
50	439706	AW872527	Hs.59761	ESTs, Weakly similar to DAP1_HUMAN DEATH	hi-lo-lo-hi hi-lo-lo-hi
-	450088	AW292933	Hs.254110	ESTs	hi-lo-lo-hi
	414219	W20010	Hs.75823	ALL1-fused gene from chromosome 1q	hi-lo-lo-hi
	419201	M22324	Hs.1239	alanyl (membrane) aminopeptidase (aminop	hi-lo-lo-hi
	426263	Al908774	Hs.259785	carnitine palmitoyitransferase I, liver	hi-lo-lo-hi
55	456236	AF045229	Hs.82280	regulator of G-protein signalling 10	hi-lo-lo-hi
	456607	A1660190	Hs.106070	cyclin-dependent kinase inhibitor 1C (p5	hi-lo-lo-hi
	408437	AW957744	Hs.278469	lacrimal proline rich protein	hi-lo-lo-hi
	421180	BE410992	Hs.258730	heme-regulated initiation factor 2-alpha	hi-lo-lo-hi
60	413437	BE313164	Hs.75361	gene from NF2/meningioma region of 22q12	hi-lo-lo-hi
00	432415	T16971	Hs.289014	ESTs, Weakly similar to A43932 mucin 2 p	hi-lo-lo-hi
	449230 417979	BE613348	Hs.211579	melanoma cell adhesion molecule GTP cyclohydrolase I feedback regulatory	hi-lo-lo-hi
	421877	AU077284 AW250380	Hs.83081 Hs.109059	mitochondrial ribosomal protein L12	hi-lo-lo-hi hi-lo-lo-hi
	412482	A1499930	Hs.334885	mitochondrial GTP binding protein	hi-lo-lo-hi
65	428423	AU076517	Hs.184276	solute carrier family 9 (sodium/hydrogen	hi-lo-lo-hi
	422947	AA306782	Hs.122552	G-2 and S-phase expressed 1	hi-lo-lo-hi
	441072	AW275480	Hs.39504	hypothetical protein MGC4308	hi-lo-lo-hi
	415938	BE383507	Hs.78921	A kinase (PRKA) anchor protein 1	hi-lo-lo-hi
<b>-</b> 0	432278	AL137506	Hs.274256	hypothetical protein FLJ23563	hì-lo-lo-hi
70	446651	AA393907	Hs.97179	ESTs	hi-lo-lo-hi
	431515	NM_012152		endothelial differentiation, lysophospha	hi-lo-lo-hi
	445345	AW003850	Hs.12532	chromosome 1 open reading frame 21	hi-lo-lo-hi
	458965	AA010319	Hs.60389	ESTs	hi-lo-lo-hi
75	438321	AA576635	Hs.6153	CGI-48 protein	hi-lo-lo-hi
13	416783	AA206186	Hs.79889	monocyte to macrophage differentiation-a	hi-lo-lo-hi
	453563 432303	AW608906	Hs.181163	hypothetical protein MGC5629	hì-lo-lo-hi
	432393 433914	AW205863 AF108138	Hs.133988 Hs.112160	hypothetical protein FKSG28 Homo sapiens DNA helicase homolog (PIF1)	hi-lo-lo-hi hi-lo-lo-hi
	433914	X90725	Hs.77597	polo (Drosophia)-like kinase	ni-to-to-ni hi-lo-lo-hi
80	432375	BE536069	Hs.2962	S100 calcium-binding protein P	hi-lo-lo-hi
	440773	AA352702	Hs.37747	Homo sapiens, Similar to RIKEN cDNA 2700	hi-lo-lo-hi
	415994	NM_002923		regulator of G-protein signalling 2, 24k	hi-lo-lo-hi
				174	

	412722	AI343300	Hs.15091	ESTs	hi-lo-lo-hi
	446839	BE091926	Hs.16244	mitotic spindle coiled-coil related prot	hi-lo-ko-hi
		NM_000346		SRY (sex determining region Y)-box 9 (ca	hì-lo-lo-hi
	428862 439108	AW163034	Hs.6467	synaplogyrin 3	hi-lo-lo-hi
5		AW449612		ESTs	hi-lo-lo-hi
J	430178 421733		Hs.152475	fibroblast growth factor receptor 3 (ach	hi-lo-lo-hi
		AL119671 AL133619	Hs.1420	Homo sapiens mRNA; cDNA DKFZp434E2321 (f	hi-lo-lo-hi
	452410		11- 024440	hypothetical protein FLJ20647	hi-lo-lo-hi
	430132	AA204686 AA236291	Hs.234149		hi-lo-lo-hi
10-	428297		Hs.183583	serine (or cysteine) proteinase inhibito	
IV	413142	M81740	Hs.75212 Hs.174070	ornithine decarboxylase 1 ubiquitin carrier protein	hi-lo-lo-hi hi-lo-lo-hi
	427239	BE270447			11-10-10-111 hi-to-to-hi
	409738	BE222975	Hs.56205	insulin induced gene 1 chromosome 1 open reading frame 21	ni-to-to-ni hi-to-to-hi
	410748	BE383816	Hs.12532		hi-lo-to-hi
15	424506	AF220490	Hs.149623	group III secreted phospholipase A2	hi-lo-lo-hi
13	447333	BE090580	Hs.70704	hypothetical protein dJ61688.3	hi-lo-lo-hi
	414761	AU077228	Hs.77256	enhancer of zeste (Drosophila) homolog 2	hi-lo-lo-hi
	419602	AW248434	Hs.91521	hypothetical protein	hi-lo-lo-hi
	411669	BE612676	Hs.303116	stromal cell-derived factor 2-like 1	nH0-10-m hi-lo-lo-hi
20	452322	BE566343	Hs.28988	glutaredoxin (thioltransferase)	hi-lo-lo-hi
20	426006	R49031	Hs.22627	ESTs	
	457465	AW301344	Hs.122908	DNA replication factor	hi-lo-lo-hi hi-lo-lo-hi
	406867	AA157857	Hs.182265	keratin 19	hi-lo-lo-hi
	407230	AA157857	Hs.182265	keratin 19	hi-lo-lo-hi
25	446681	AJ003624	Hs.15896	kendrin phosphoglycerate mutase 2 (muscle)	hi-lo-lo-hi
25	408493	BE206854	Hs.46039	GDP-mannose 4,6-dehydratase	hi-lo-lo-hi
	439186	A1697274	Hs.105435		' hi-lo-lo-hi
	424544	M88700	Hs.150403 Hs.5794	dopa decarboxylase (aromatic L-amino aci ESTs, Wealdy similar to 2109260A B cell	hi-lo-lo-hi
	431325	AW026751 D00723	Hs.77631	glycine cleavage system protein H (amino	hi-lo-lo-hi
30	414922 438291	BE514605	Hs.289092	Homo sapiens cDNA: FLJ22380 fis, clone H	hi-lo-lo-hi
50			NS.203032	M-phase phosphoprotein 9	hi-lo-lo-hi
	418574	N28754 AU077058	Hs.54089	BRCA1 associated RING domain 1	hi-lo-lo-hi
	409342 432734	AA837396	Hs.263925	US1-interacting protein NUDE1, rat homo	hi-lo-lo-hi
			Hs.5054	CGI-133 protein	hi-lo-lo-hi
35	436087 420309	BE300296 AW043637	Hs.21766	ESTs, Weakly similar to ALU5_HUMAN ALU S	hi-lo-lo-hi
55	420309	AW043037 Al418609	Hs.71040	hypothetical protein FLJ20425	hi-lo-lo-hi
	424381	AA285249	Hs.146329	protein kinase Chk2	hi-lo-lo-hi
	442547	AA306997	Hs.217484	ESTs, Weakly similar to ALU1_HUMAN ALU S	hi-lo-lo-hi
	430376	AW292053	Hs.12532	chromosome 1 open reading frame 21	hi-lo-lo-hi
40	434666	AF151103	Hs.112259	T cell receptor gamma locus	hi-to-lo-hi
<del>+</del> 0	412330	NM 005100		A kinase (PRKA) anchor prolein (gravin)	hi-lo-lo-hi
	452123	Al267615	Hs.38022	ESTs	hi-lo-lo-hi
	424893	AW295112	Hs.153648	Homo sapiens cDNA FLJ13303 fis, clone OV	hì-lo-lo-hi
	428057	Al343641	Hs.185798	ESTs	hì-lo-lo-hì
45	431566	AF176012	Hs.260720	J domain containing protein 1	hi-lo-lo-hi
13	439979	AW600291	Hs.6823	hypothetical protein FLJ10430	hi-lo-lo-hi
	418836	AI655499	Hs.161712	ESTs	hi-lo-lo-hi
	433757	Al949974	Hs.152670	ESTs	hi-lo-lo-hi
	425236	AW067800	Hs.155223	stanniocatcin 2	hi-lo-lo-hi
50	426215	AW963419	Hs.155223	stanniocalcin 2	hi-lo-lo-hi

## TABLE 2B

Pkey: Unique Eos probeset identifier number CAT number: Gene cluster number Accession: Genbank accession numbers

Piege   CAT Number   Accession   109784-1	,	ACCESSIO	HI. GEHDAIK ALL	ession numbers
1075961   ACCESTED AND SERVE AND S		Pkev	CAT Number	Accession
### 498051   109869_1   AM09012 AM7518 AM08303 AM76904 AM07093 AM07093 AM07093   ### 19805   109861_1   AM09012 AM7518 AM08303 AM76905 AM07093 AM07093   ### 19805   109861_1   AM09012 AM7518 AM08303 AM76905 AM07093 AM76905   ### 19805   109861_1   AM09012 AM76905 AM07093 AM07093 AM76905 AM7690				1.4
ANGESES AMOTHER PARTIES AMOSESS AMOSES				
499123   11014_1   A0053403 ANDRESS	10	405001	103035_1	
19216   194664   19501   194111   19511   194111   19511   1	10	409123	110143 1	
140451   120416_1				
14048 12081   A. A. A. SECTION AND SECTIONS OF CHILD STANDARD CONTROL				
155 411033 123046.1 411031 123046.1 411031 123046.1 141031 14103				
## # # # # # # # # # # # # # # # # # #	15			
## # # # # # # # # # # # # # # # # # #	13			
4 11701   1254465   162400   1 AWSSSSS AMSSSS AMSSS AMSSSS AMSSS AMSSSS AMSSS AMSSSS A				
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4 12419 12394-1 1 12394-1				
A	20			
412492   130802,1   AMPSEB04 AA368539 AA112267	20	412419	1253410_1	
413567   131807   1   1807   1   1807   1   18080   1   1808081		440400	420002 4	
143581   133580.1				
41599 1374313   1 41596 153573   1 41596				· · · · · · · · · · · · · · · · · · ·
## # # # # # # # # # # # # # # # # # #	25			
415308   153372_1   FI0230T R1748 Z40028 H14747   FI0251 R1749 Z4305 H2070	23			
415516   153918_1   F11411 F15277 2/3915 H20760     41651   150519_1   41651   150519_1   41651   150519_1     41651   150519_1   41651   150519_1   41651   150519_1   41651   150519_1   41651   150519_1   41651   150519_1   41651   150519_1   41651   150519_1   41651   150519_1   41651   416519_1   41651				
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41654 1605019_1   41654 163427_1   41654 163427_1   41654 163427_1   41654 163427_1   416556 163647_1   416236 17184_1   4165666_1   416236 17184_1   416236 1  416236 17184_1   416236 1  416236				
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41829				
148574   1789_T   1				
Main	25			
41955	33	418574	17690_1	N28754 N28747 Al568146 Al979339 AA322671 AA322672 AW955043 Al990326 AA776406 Al016250 AA843678 AW451882 N23137 N23129
42081   19877_1				
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42476 22861_1 AL036533 F11794 F11783 H18027 E16008 H2927 R19493 AW134660 Al289437 AL 133995 AA057405 N78357 AA917450 Al002692 T09262 T05008 H28028 A20306.1 4 A23021 AA034729 AA034739 AA0344370 AA03422 AA03424 AA034		422128	211994_1	AW881145 AA490718 M85637 AA304575 T06067 AA331991
42476 22861_1 AL036533 F11794 F11783 H18027 E16008 H2927 R19493 AW134660 Al289437 AL 133995 AA057405 N78357 AA917450 Al002692 T09262 T05008 H28028 A20306.1 4 A23021 AA034729 AA034739 AA0344370 AA03422 AA03424 AA034		423028	224062_1	H90946 AA320597 AW954970 BE143680
45				AL035633 F11794 F11783 H18042 T66089 H29379 R19493 AW134660 Al299437 AL133995 AA057405 N78357 AA917450 Al002692 T09262
42885 233006.1 AA33221 5AA403110 AW956299 42843 24194.1 AA34572 AA47080 H97831 AA350358 BE166712 42890 258778.1 AA35627 AW062361 AW813419 AW816041 AI744949 42891 289818.1 AA35627 AW70899 AA694295 428161 28785.1 AA36851 AA47089 AA694288 428161 28785.1 AA36851 AA47089 AA694288 428161 28785.1 AA36851 AA47089 AA694288 428161 28785.1 AA684768 AW978427 IAA52275 AA447312 42963 30568.1 AA684768 AW978427 IAA52275 AA447312 42963 30568.1 AA684768 AW978427 IAA52275 AA447312 43068 312849.1 AA684768 AW978427 IAA52275 AA447312 43043 31808.1 AA68468 AW6505 AA947668 43043 31808.1 AA64695 AW897144 AL122069 AW39292 AI988826 43183 337824.1 AA516420 C14818 C14815 C15161 C15068 D80763 D60656 AW970134 AA543007 D81004 D60184 AI488371 D60382 D60181 C15876 437307 34114.1 AA972747 AA525237 AA846376 AA470580 AA676538 BE27147 AW291971 AA017125 AI198417 AI365213 AI16842 AI337018 433075 35820.1 AM6566 AA58869 AA686876 AA476580 AA476328 AA46826 AA58869 AA686876 AA676602 432340 345240.1 AA54622 AA68438 AA686876 AA476580 AA76538 BE27147 AW291971 AA017125 AI198417 AI365213 AI16842 AI337018 AN566 AA58689 AA66818 AA6868 AA688800 AA418325 AA418378 NT1981 AL103634 AA428301 AA488275 AA232975 AL036861 BE277220 BE387505 N99710 AW375004 AA46286 AA58889 AA686876 AA76023 BE27147 AW291971 AA017125 AI198417 AI365213 AI16842 AI337018 AA6868 AA6868 AA6868 AA6868 AA68680 AA418325 AA418378 NT1981 AL103634 AA428301 AA488275 AA232975 AL036861 BE277220 BE387505 N99710 AW375004 AA418285 AL079661 HE3776 AA68686 AA6868 AA686			-	
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N99710 AW375004 AA418268 AL079651 H85743 AW902319 AW805907 AA984366 T92310 AA405425 AA421732 Al656841 AW300968		100010	00020	
AW593418 T92267 BE464032 AW473548 Al359502 BE552306 Al990196 AW518351 Al239559 AW590963 AA018359 Al273737 AL042658 AA411308 AA402810 H38111 AlW013931 AW3664824 AW3762424 SAW376124 Al290202 Al292121 AA340647 BE613672 BE409874 AA351915 BE617026 BE019588 AW402692 AW247466 R59223 AA134761 BE254019 BE265105 D63316 BE313080 BE547713 BE536578 BE546749 AA324165 H17386 BE253377 R87598 H29072 AA350980 BE076629 BE253957 AA532613 BE252486 AW804459 D30966 R87959 AA091832 BE005398 AA628622 AA994155 R76593 AF147390 R76594 A36716 425440_1 A436862 42814_2 A436862 42814_2 A436862 42814_2 A436862 42814_2 A436869 46651_1 A38882 466649_1 A38882 466649_1 A438882 466649_1 A438884 47806_1 A39884 467544_1 A499086 A68133_1 AA947354 AA829660 Al687296 A39988 477806_1 A99086 468133_1 AA947354 AA82960 Al687296 A39988 477806_1 A4886167 F21558 F31418 F35624				
AA411308 AA402810 H38111 AW013931 AW366432 AW752435 AW376124 AI292020 AI292121 AA340647 BE613672 BE409874 AA351915 BE617026 BE019588 AW402692 AW247466 R59233 AA134761 BE256105 D83316 BE313080 BE547713 BE536578 BE546749 AA324185 H17386 BE253377 R87598 H29072 AA350980 BE076629 BE253957 AA532613 BE252486 AW804459 D30966 R87959 AA091832 BE005398 AA628622 AA994155 R76593 AF147390 R76594 A39023 398093_1 AI692552 AI393343 AI800510 AI377711 F24263 AA661876 A36716 425440_1 A433540 AA728984 AA804981 A36862 42814_2 A8829_1 BE514383 AA071273 AW247987 AW673286 BE312102 AW749824 BE071985 AW577383 BE071945 BE072005 AW577355 BE071965 AW239231 BE072000 BE071960 AW577360 AW749830 AW373020 X97303 AW999522 BE000192 BE562219 BE266655 BE264970 A38869 46651_1 AF075009 R63109 R63068 A38889 467544_1 AW502384 AI982587 AA828822 A39888 477806_1 AW502384 AI982587 AA828688 A480151 487109_1 AA868167 F21558 F31418 F35624	65			
### BE617026 BE019588 AW402692 AW247466 R59233 AA134761 BE256105 D63316 BE313080 BE547713 BE536578 BE546749  ### AA324185 H17386 BE253377 R87598 H29072 AA350980 BE076629 BE253957 AA532613 BE252486 AW804459 D30966 R87959 AA091832  ### BE005398 AA628622 AA994155  ### BE005398 AA628622 AA994155  ### BE005398 AA628622 AA994155  ### R76593 AF147390 R76594  ### A1692552 A1393343 A1800510 A1377711 F24263 AA661876  ### A36716 425440_1 A433540 AA728984 AA804981  ### A36862 42814_2 A1821940 N67106 A1744264 AA808846 AA643417 AA643416 Z70715  ### BE514383 AA071273 AW247987 AW673286 BE312102 AW749824 BE071985 AW577383 BE071945 BE072005 AW577355 BE071965 AW239231  ### BE514383 AA071273 AW247987 AW673286 BE312102 AW749824 BE071985 AW577383 BE071945 BE072005 AW577355 BE071965 AW239231  ### BE514383 AA071273 AW247987 AW673286 BE312102 AW749824 BE071985 AW577383 BE071945 BE072005 AW577355 BE071965 AW239231  ### BE514383 AA071273 AW247987 AW673286 BE312102 AW749824 BE071985 AW577383 BE071945 BE072005 AW577355 BE071965 AW239231  ### BE514383 AA071273 AW247987 AW673286 BE312102 AW749824 BE071985 AW577383 BE071945 BE072005 AW577355 BE071965 AW239231  ### BE514383 AA071273 AW247987 AW673286 BE312102 AW749824 BE071985 AW577383 BE071945 BE072005 AW577355 BE071965 AW239231  ### BE514383 AA071273 AW247987 AW673286 BE312102 AW749824 BE071985 AW577383 BE071945 BE266655 BE2664970  ### A48869 46651_1	05			
A324185 H17386 BE253377 R87598 H29072 AA350980 BE076629 BE253957 AAS32613 BE252486 AW804459 D30966 R87959 AA091832  434280 382816_1 BE005398 AA628622 AA994155 R76593 AA628622 AA994155 A36903 398093_1 A692552 A1393343 A1800510 A1377711 F24263 AA661876 A36716 425440_1 A433540 AA728984 AA804981 A36862 42814_2 A1821940 N67106 A1744264 AA808846 AA643417 AA643416 Z70715 BE514383 AA071273 AW247987 AW673286 BE312102 AW749824 BE071985 AW577383 BE071945 BE072005 AW577355 BE071965 AW239231 BE072000 BE071960 AW577360 AW749830 AW373020 X97303 AW999522 BE000192 BE562219 BE266655 BE264970 A7675009 R63109 R63068 A38882 466649_1 AA827695 AA833754 AW978946 A39888 477806_1 AW502384 A1982587 AA828622 A39848 477806_1 AA947354 AA829660 A1687296 A39848 477806_1 AW579249 D63277 AA846988 A40151 487109_1 AA868167 F21558 F31418 F35624				
70				
70		434300	202016 1	
435023 398093_1	70			
75 436716 425440_1 436862 42814_2 437576 43892_1 82514383 A071273 A0728984 AA804981 A651_1 AF075009 R63109 R63068 438882 466649_1 438880 467544_1 439846 468133_1 80 43946 468133_1 43946 468133_1 43946 468133_1 43948 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 43988 477806_1 439948 487109_1 438698 440151 487109_1 AA868167 F21558 F31418 F35624	70			
75				
75 438757 43892_1 BE514383 AA071273 AW247987 AW673286 BE312102 AW749824 BE071985 AW577383 BE071945 BE072005 AW577355 BE071965 AW239231 BE072000 BE071960 AW577360 AW749830 AW373020 X97303 AW999522 BE000192 BE562219 BE266655 BE264970  438869 46651_1 AF075009 R63109 R63068  438980 467544_1 AA827695 AA833754 AW978946  438980 467544_1 AW502384 AI982587 AA828822  43946 468133_1 AA947354 AW502384 AI982587 AA82882  43948 477806_1 AW979249 D63277 AA846968  440151 487109_1 AA868167 F21558 F31418 F35624				
75				
438869 46651_1	75	43/5/6	43892_1	
438882 466649_1	13	400000	10054 4	
438980 467544_1 AW502384 Al982587 AA828822 439046 468133_1 AA947354 AA829660 Al687296 439848 477806_1 AW979249 D63277 AA846968 440151 487109_1 AA868167 F21558 F31418 F35624				
439046 468133_1				
80 439848 477806_1 AW979249 D63277 AA846968 440151 487109_1 AA868167 F21558 F31418 F35624				
440151 487109_1 AA868167 F21558 F31418 F35624	90			
	δU			
44050/ 4956//_1 H06994 BE147898				
		440507	495677_1	HU6994 BE147898

5		59994_1 600667_1	AA973905 Al299888 AA917019 H63235 T90771 AA974603 Al984319 AW340495 Al038316 Al344631 Al261653 AA262496 AV648929 AA305356 D61644 D78724 Al140497 AW749625 AW749626 AW749644 AV655234 AW966332 AA340239 BE090517 AW970792 AW264490 AW014985 F27436 AA947336 F15843 H89338 AA563626 F17712 BE546579 AA421821 AA284852 AA477751 AW025245
10	448150 448489 448631	722246_1 752165_1 765247_1 772996_1 77790_1	AW025243 BE244285 C18429 H42373 AI820706 AI379786 R55439 AW276142 AI472167 AI990315 R32175 AI523875 R45782 R45781 AI554923 AI902356 BE614081 W01988 AW500790
15		9163_1	AL133619 AA468118 AA383064 Al476447 T09430 Al673758 AA524895 Al581345 Al300820 AW498812 AA256162 Al559724 Al685732 AA602400 AA905453 Al204595 AW166541 AA157456 AA156269 AA383652 AA431072 AW592707 Al435410 AW272464 Al215594 AA622747 R74039 N35031 Al804128 AW513621 AA668351 Al026826 Al493388 AA614641 W81604 Al567080 Al214351 AA730140 Al125754 Al200813 Al269603 Al565082 Al807095 Al476629 AA505999 Al368449 Al686077 Al582930 AW085038 AA757863 AA730154 Al767072 AA468316 Al734130 Al734138 AA426284 AA433997 Al741241 AW043563 Al732741 Al732734 AA437369 AA425820 AA664048 R74130
20	452654 454775 455019	918078_1 925931_1 1234106_1 1249138_1	BE144022 BE143969 BE143915 BE004783 BE004947 Al911790 BE160229 AW819879 AW820179 AW819882 AW819876 AW820169 BE153201 AW993736 BE152911 AW850818 AW850833 AW851100
25	455619 455653 455729 455824	1271871_1 1346387_1 1348742_1 1353792_1 1372880_1	BE148152 BE148133 BE148159 BE148132 AW885107 BE063853 BE063955 BE063866 BE063705 BE063846 BE061416 BE063844 BE154075 BE153973 BE064861 BE153852 BE153847 BE064684 BE153602 BE065075 BE154018 BE064772 BE064842 BE153557 BE153509 BE072092 BE072106 BE072086 BE072098 BE072103 BE143703 BE143631 BE143629 BE143702
30	455956 456123 457133	1387163_1 1534442_1 29066_1	BE162704 BE162705 BE162732 BE162702 BE162694 R00602 Z42921 F06132 M54968 NM_004985 Al808924 AL135130 AW242010 AA476848 AI740449 M17087 K03210 M35505 M35504 L00049 Al186585 W35273 X01669 X02825 W23635 Al554920 Al539465 AA425263 Al469981 W21091 T28976 AW977922 BE550180 AW664973 Al148939 AW117295 AA811229 Al343010 AA766141 BE219368 N95249 AA280396 AW504574 AA232870 AI770018 AA262948 AW450230 AW362890 AW609417 AW499941
35			AA425857 AW380665 AA830647 AA282180 T27356 H85307 AA861543 AA356548 AA356410 AW860656 AW860647 AW938103 AW860649 AI567016 N70374 AW474707 AA505084 AA082195 AW949515 AA361728 N33863 AA411821 AA401640 AW994461 AL120766 AI500024 AW771891 H84567 D51551 AA330600 R14184 AI301629 N64676 AV569669 AI69760 AI004579 AA287927 AW453052 AW601642 AA676681 AA737010 AA872481 AA281094 AA564243 BE464958 BE049265 AW167917 AA843916 AA525301 AI015987 N25230 AI889481 AW173466 AA937541 AI334416 AI676214 AI281159 AA553559 AA582189 AA255527 AW160515 AA670007 H08199 AA808271 AA281015 W47527 AA649252
40			AI364302 AA889246 R40473 H02312 AA648116 AA342730 AA243624 R99351 R41588 R49696 AA854442 F01713 AA213685 AA721296 R79833 H84241 R70668 H85554 AA223758 N95349 AI374913 AI306683 AA015609 AA918548 AI453570 AA772321 AI692775 AA195733 AI474563 AW873048 AI209133 AI028182 AI374920 AW572807 AA406223 AA833684 T97255 H69138 AA36906 AW119162 N31974 AI890548 N39418 AA864877 AA679469 BE350651 N41020 AI050915 F00075 AA864878 N26970 AA828898 AW019991 AW796631 AW993262 N48532 BE564662 AV654063 AI754461 AW945712 C03289 AV655314 AV659070 AV659808 AV660435 H70113 C05323 R91984 H96949 AV658936 AV658879
45			H69137 AA384411 AA412584 C02749 W32014 R58168 C05526 BE536017 N24354 AA287991 N80109 F05452 R12740 H08297 AL138354 AW020801 BE178443 BE178018 BE178336 BE178360 BE178107 BE178385 BE178215 BE17816 BE178147 BE178352 BE178422 BE178424 BE178043 BE178093 BE178460 BE178356 BE178441 BE178438 BE178467 Al091259 BE177839 BE178094 R28455 BE177844 BE178100 AA262387 R70669 W80934 W93668 A4256711 BE178141 BE177893 BE178444 AA167718 H69694 BE178017 BE178029 BE177999 BE177936 AA095144 N32462 AA281203 AA281183 W47526 W05015 R34165 R35396 T97366 R79640 W25258 R99450 AW368425 BE178196 R26447
50	457952 458956	44256_1 83645_1	C03146 C03683 U25750 AI792472 AA487379 AI872282 AA487262 R22383 AI865750 R21832 AA593628 AW571869 AA377191 R78814 T27193 BE220675 AA345621 AA009992

## TABLE 2C

Pkey: Unique number corresponding to an Eos probeset
Ref: Sequence source. The 7 digit numbers in this column are Genbank Identifier (GI) numbers. "Dunham I. et al." refers to the publication entitled "The DNA sequence of human chromosome 22." Dunham I. et al. (1999) Nature 402:489-495.
Strand: Indicates DNA strand from which exons were predicted.
NL position: Indicates nucleotide positions of predicted exons. 5

10	Pkey	Ref	Strand	Nt_position
10	400481	8439853	Plus	112433-112541
	400501	9796227	Minus	12479-12619
	400713		Minus	43185-43394
	400769	8131628	Plus	28671-29795
15	400818	8569994	Plus	172644-172765,173085-173200
15	400881	2842777	Minus	91446-91603,92123-92265
	400882		Minus	110431-110708
	400965	7770576	Minus	173043-173564
	400986	8085497	Minus	63140-63319
20	400995	8099094	Plus	141186-141601
20	401093	8516137	Minus	22335-23166
	401178	9438616	Minus	133663-133812
	401192	9719502	Minus	69559-70101
	401209	7712287	Plus	164932-165112
25	401405	7768126	Minus	69276-69452,69548-69958
25	401416	7452889	Minus	121456-121626
	401419	7452889	Minus	136389-136508
	401444	8346725	Plus	90895-90994,93070-93213
	401512	7622346	Plus	136399-136557
20	401563	8247910	Plus	91395-91763
30	401600	4388746	Minus	27363-27518,28727-28891,29526-29731
	401750	9828651	Plus	82143-82270,89284-89373,90596-90770,95822-96001,96688-96775,96870-96992,98046-98138
	401757	7239630	Plus	88641-88751
	401839	7656637		1016-1086,2751-2967,3241-3348,26677-26831
35	401849	7770425	Plus	129375-129483,129597-129720
33	401952	3319121	Minus	53770-53979
	401966	3126781	Plus	29397-29918
	402082		Minus	190046-190183
	402101	8117697	Plus	134308-134487,135402-135587,136421-136548
40	402106	8131652	Plus	3717-3848
+0	402163	8568936	Plus	166996-167119
	402185	8576002	Plus	25486-25639
	402240 402249	7690131 7704953	Plus	104382-104527,106136-106372
	402249	8099267	Minus	107636-107813,108694-108824,110435-110502,113182-113386
45	402396	1905896	Minus Plus	13714-15440 4426-4648
73	402356		Minus	
	402532	9800951	Minus	71266-72351 180240-180558
	402559	9864273	Plus	33539-33715
	402575	9884830	Minus	109742-109883
50	402602	7239666	Plus	6785-6972,7478-7575
	402758	9213869	Plus	87638-87924
	402786	9715046	Plus	47624-47795
	402807	6456148	Minus	101542-101660,103476-103656
	402810	6010110	Plus	12715-12856,13527-13643
55	402964	9581599	Minus	46624-46784
	403046	3540153	Minus	55707-55859,56369-56511
	403055	8748904	Minus	109532-110225
	403217	7630969	Plus	54089-54163,55427-55623
	403218	7630969	Plus	58039-58149
60 -	403291	7230870	Plus	95177-95435
	403328	8469086	Minus	120428-120703
	403654	8736093	Minus	28634-28758
	403704	4982546	Minus	8850-8996
	403708	5705981	Minus	134394-134812
65	403725	7534031	Plus	86737-86843
	403739	7630882	Plus	44563-44766,48209-48483,52255-52495
	403740	7630882	Plus	86504-87227
	403745	7652036	Minus	67610-68002
<b>-</b> 0	403746	7652036	Plus	93612-93887
70	403885	7710403	Minus	53259-53524
	403893	7710581	Minus	5435-7846
	403947	7711923	Plus	38657-38817
	404039	8698763	Plus	81889-82011
75	404054	3548785	Plus	66713-69175
75	404058	3548785	Plus	99397-101808
	404108	8247074	Minus	63503-64942
	404211	5006246	Plus	185728-185885,194575-194686
	404277	1834458	Minus	91665-91946
90	404384	8887028	Minus	38055-38156,42175-42391,43435-43553
80	404407	7329316	Minus	48154-48499
	404489	8113772	Plus	98183-98480
	404527	8152087	Plus	127737-127796,128080-128210,129888-130054,132545-132869

	404528	8152087	Plus	135325-135486
	404661	9797073	Plus	33374-33675,33769-34008
	404663	9797133	Plus	29885-30514
_	404956	7387343	Pius	55883-56203
5	405011	6139150	Plus	117359-117612
	405044	7596797	Minus	98903-101141
	405102	8076881	Minus	120922-121296
	405109	8096886	Minus	30301-30518
	405188	6649489	Plus	134573-134678
10	405231	7249032	Minus	109793-109969
	405365	2275192	Minus	119867-120372,120481-120824,121029-121357
	405387	6587915	Minus	3769-3833,5708-5895
	405396	6624129	Minus	89965-90273
	405429	7321905	Minus	51577-51723
15	405435	7408068	Minus	51704-51841,53581-53767
	405446	7582529	Plus	99136-99313
	405503	9211311	Minus	51198-51314
	405525	9558552	Minus	19699-19828
_	405529	9581957	Minus	38944-39213
20	405610	5757553	Minus	71907-72080
	405802	5924004	Minus	27743-28264
	405811	4902753	Plus	5128-5248
	406180	7283201	Minus	38923-39107
	406207	5923650	Minus	162607-162800
25	406302	8575868	Plus	168961-169150,169610-169769

Table 3A shows the Seq ID No, Pkey, ExAcon, UnigeneID, and Unigene Title for all of the sequences in Table 4.

Pkey: Unique Eos probeset identifier number
ExAccn: Exemplar Accession number, Genbank accession number
UnigenelD: Unigene number
Unigene Title: Unigene gene title
Seq ID No: Seq ID number correlation for those sequences in Table 4

1.0	Pkey	ExAccn	UnigenelD	Unigene Title	Seq ID No
10	415539	AJ733881	Hs.72472	BMP-R1B	Seq ID No 1 & 2
	448988	Y09763	Hs.22785	gamma-aminobutyric acid (GABA) A recepto	Seq ID No 3-10
	403740			NM_001076*:Homo sapiens UDP glycosyltran	Seq ID No 11 & 12
	408633	AW963372	Hs.46677	PRO2000 protein	Seq ID No 13 & 14
	408660	AA525775		ESTs, Moderately similar to PC4259 ferri	Seq ID No 15 & 16
15	409051	AA080912		gb:zn04d03.r1 Stratagene hNT neuron (937	Seq ID No 17
	409123	AA063403		gb:zm04d12.s1 Stratagene comeal stroma	Seq ID No 18
	415787	H01463	Hs.93534	ESTs	Seg ID No 19-21
	415999	AA172179	Hs.294029	ESTs	Seq ID No 22
	416225	AA577730	Hs.188684	ESTs, Weakly similar to PC4259 ferritin	Seg ID No 23
20	420757	X78592	Hs.99915	androgen receptor (dihydrotestosterone r	Seq ID No 24 & 25
	429163	AA884766		gb;am20a10.s1 Soares_NFL_T_GBC_S1 Homo s	Seq ID No 26
	429441	AJ224172	Hs.204096	lipophilin B (uteroglobin family member)	Seq ID No 27 & 28
	431099	Y13367	Hs.249235	phosphoinositide-3-kinase, class 2, alph	Seq ID No 29 & 30
	432432	AA541323	Hs.115831	ESTs	Sea ID No 31
25	432435	BE218886	Hs.282070	ESTs	Seg ID No 32 & 33
23	432527	AW975028	Hs.102754	ESTs	Seg ID No 34
	435876	AW612586	Hs.160271	G protein-coupled receptor 48	Seq ID No 35 & 36
	438233	W52448	Hs.56147	ESTs	Seq ID No 37-40
		AW602166	Hs.222399	CEGP1 protein	Seg ID No 41 & 42
30	439569			ESTs	Seq ID No 43
30	440819	A1809444	Hs.202108		Seq ID No 44
	442832	AW206560	Hs.253569	ESTs	Seg ID No 45 & 46
	447342	A)199268	Hs.19322	Homo sapiens, Similar to RIKEN cDNA 2010	
	447499	AW262580	Hs.147674	protocadherin beta 16	Seq ID No 47 & 48
25	451411	AA017492	Hs.135655	EST	Seq ID No 49
35	451720	AW970985	Hs.290853	ESTs	Seq ID No 50 & 51

Table 3B shows the accession numbers for those Pkey's lacking UnigeneiD's for table 3A. For each probeset is listed gene cluster number from which oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

5	Pkey	CAT Number	Accession
	408660	107294_1	AA525775 AA056342 AI538978 AW975281 AA664986
	409051	109699_1	AA080912 AA075318 AA083403 AA076594 AA078992 AA084926 AA081881 AA113913 AA113892 AA083821 AA134801 AA082953 AA070343
			AA062835 AA075419 AA063293 AA071252 AA078900 AA062836 AW974305
	409123	110143_1	AA063403 AA070823 AA070050
10	429163	300543_1	AA884766 AW974271 AA592975 AA447312

Table 3C shows genomic positioning for those Pkey's lacking Unigene ID's and accession numbers in table 3A. For each predicted exon is listed genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed.

 Pkey
 Ref
 Strand
 Nt\_position

 5
 403740
 7630882
 Plus
 86504-87227

Table 4:

Seq ID NO: 1 <u>DNA sequence</u> Nucleic Acid Accession #: NM\_001203 5 Coding sequence: 274..1782 CGCGGGGCGC GGAGTCGGCG GGGCCTCGCG GGACGCGGGC AGTGCGGAGA CCGCGGCGCT 10 GAGGACGCGG GAGCCGGGAG CGCACGCGCG GGGTGGAGTT CAGCCTACTC TTTCTTAGAT 120 GTGAAAGGAA AGGAAGATCA TTTCATGCCT TGTTGATAAA GGTTCAGACT TCTGCTGATT 180 CATAACCATT TGGCTCTGAG CTATGACAAG AGAGGAAACA AAAAGTTAAA CTTACAAGCC 240 TGCCATAAGT GAGAAGCAAA CTTCCTTGAT AACATGCTTT TGCGAAGTGC AGGAAAATTA 300 AATGTGGGCA CCAAGAAAGA GGATGGTGAG AGTACAGCCC CCACCCCCCG TCCAAAGGTC 360 15 TTGCGTTGTA AATGCCACCA CCATTGTCCA GAAGACTCAG TCAACAATAT TTGCAGCACA 420 GACGGATATT GTTTCACGAT GATAGAAGAG GATGACTCTG GGTTGCCTGT GGTCACTTCT 480 GGTTGCCTAG GACTAGAAGG CTCAGATTTT CAGTGTCGGG ACACTCCCAT TCCTCATCAA AGAAGATCAA TTGAATGCTG CACAGAAAGG AACGAATGTA ATAAAGACCT ACACCCTACA CTGCCTCCAT TGAAAAACAG AGATTTTGTT GATGGACCTA TACACCACAG GGCTTTACTT ATATCTGTGA CTGTCTGTAG TTTGCTCTTG GTCCTTATCA TATTATTTTG TTACTTCCGG 660 20 720 TATAAAAGAC AAGAAACCAG ACCTCGATAC AGCATTGGGT TAGAACAGGA TGAAACTTAC 780 ATTCCTCCTG GAGAATCCCT GAGAGACTTA ATTGAGCAGT CTCAGAGCTC AGGAAGTGGA 840 TCAGGCCTCC CTCTGCTGGT CCAAAGGACT ATAGCTAAGC AGATTCAGAT GGTGAAACAG 900 ATTGGAAAAG GTCGCTATGG GGAAGTTTGG ATGGGAAAGT GGCGTGGCGA AAAGGTAGCT GTGAAAGTGT TCTTCACCAC AGAGGAAGCC AGCTGGTTCA GAGAGACAGA AATATATCAG 960 25 ACAGTETTEA TEAGGCATEA AAACATTTTE GETTTCATTE CTGCAGATAT CAAAGGGACA GGGTCCTGGA CCCAGTTGTA CCTAATCACA GACTATCATE AAAATGGTTC CCTTTATGAT 1080 1140 TATCTGAAGT CCACCACCCT AGACGCTAAA TCAATGCTGA AGTTAGCCTA CTCTTCTGTC AGTGGCTTAT GTCATTTACA CACAGAAATC TTTAGTACTC AAGGCAAACC AGCAATTGCC 1200 1260 30 CATCGAGATC TGAAAAGTAA AAACATTCTG GTGAAGAAAA ATGGAACTTG CTGTATTGCT 1320 GACCTGGGCC TGGCTGTTAA ATTTATTAGT GATACAAATG AAGTTGACAT ACCACCTAAC 1380 ACTCGAGTTG GCACCAAACG CTATATGCCT CCAGAAGTGT TGGACGAGAG CTTGAACAGA 1440 ANTCACTICC AGTOTTACAT CATGGCTGAC ATGTATAGTT TTGGCAGAGAG CTTGAACAGA
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	Cog ID NO.	8 Protein s	zemience				
	Protein Acc	cession #: 1	VP_U08822.1				
05						•	
25	1	11	21	31	41	51	
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	MEYTTOTTES	QTWNSKRTHE	HEITMPNOMV	RIYKDGKVLY	TIRMTIDAGC	SLHMLRFPMD	60
		FSYPENEMIY					120
20		RFGYVAFQNY					180
30	<b>FSRKNFPRVS</b>	YITALDFYIA	ICFVFCFCAL	LEFAVLNFLI	YNQTKAHASP	KLRHPRINSR	240
	AHARTRARSR	ACARQHQEAF	VCQIVTTEGS	DGEERPSCSA	QQPPSPGSPE	GPRSLCSKLA	300
	CCEWCKRFKK	YFCMVPDCEG	STWOOGRLCI	HVYRLDNYSR	VVFPVTFFFF	NVLYWLVCLN	360
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	Coding sequ	lence: 1309	2490				
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		a. aaaaaaa. a	amaaaaaaa a	CHCCHCCCCC	CCCRCRCCCC	つつききき かかつかかつ	
				GTGGTCGCGC			60
	TCCAAAGTTC	TTCCAGTCCT	CCTAGGCATC	TTATTGATCC	TCCAGTCGAG	AACATGTATA	120
	TCCAAAGTTC CAGAGAAGTG	TTCCAGTCCT CTCAAATCAT	CCTAGGCATC AAGTGTACAG	TTATTGATCC CTGATGAGTT	TCCAGTCGAG GTCAAAAAAT	AACATGTATA GACCACAGCG	120 180
	TCCAAAGTTC CAGAGAAGTG	TTCCAGTCCT	CCTAGGCATC AAGTGTACAG	TTATTGATCC CTGATGAGTT	TCCAGTCGAG GTCAAAAAAT	AACATGTATA GACCACAGCG	120
45	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA	TTCCAGTCCT CTCAAATCAT AGCCAAATCA	CCTAGGCATC AAGTGTACAG AGGACCCGAA	TTATTGATCC CTGATGAGTT TGTGAGCAGG	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG	AACATGTATA GACCACAGCG CCCCCTTTGT	120 180
45	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA CACTGCCTCC	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC	CCTAGGCATC AAGTGTACAG AGGACCCGAA AGCACTATCC	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA	TCCAGTCGAG GTCAAAAAT ACCTCAGAAG CACCATCGGT	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA	120 180 240 300
45	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA	CCTAGGCATC AAGTGTACAG AGGACCCGAA AGCACTATCC CATTTTTGTT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT	120 180 240 300 360
45	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT	CCTAGGCATC AAGTGTACAG AGGACCCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT	120 180 240 300 360 420
45	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA CACTGCCTCC CCITGGCAGA TTTTCTTGGC TCCTGGATGG	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG	CCTAGGCATC AAGTGTACAG AGGACCCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAAG	120 180 240 300 360 420 480
	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGCCTCCTTC	CCTAGGCATC AAGTGTACAG AGGACCCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCCTGC	TCCAGTCGAG GTCAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAAG ACGAGTATTA	120 180 240 300 360 420 480 540
45 50	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGCCTCCTTC AAAACCGCAA	CCTAGGCATC AAGTGTACAG AGGACCCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCCTGC CAATTTTCTG	TCCAGTCGAG GTCAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTCTG	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAG ACGAGTATTA GGAGAGGCCG	120 180 240 300 360 420 480 540
	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGCCTCCTTC	CCTAGGCATC AAGTGTACAG AGGACCCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCCTGC CAATTTTCTG	TCCAGTCGAG GTCAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTCTG	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAG ACGAGTATTA GGAGAGGCCG	120 180 240 300 360 420 480 540
	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAT CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AAAACGCAA TCCTCTCAGC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGGACTATCC CATTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA CCTGGCCCTC	TTATTGATCC CTGATGAGCAG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCCTGC CAATTTTCTG TGCCTGCTCC	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAA ACTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCC GTCCTCTCTG TCACTCCTGG	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGCTGG	120 180 240 300 360 420 480 540
	TCCAAAGTTC CACAGAAGTAG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGCCTCCTTC AAAACCGCAA TCCTCTCAGC TAGAGGCCAA	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG CCTTCATCCC AATATTCCCA CCTGGCCCTC GGCGACCAAC	TTATTGATCC CTGATGAGTT TOTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCCTGC CAATTTTCTG TGCCTGCT ACTAGGCAAA	TCCAGTCGAG GTCAAAAAAT ACTCCAGAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGT CCCCTCTGG CCCGGCCAGC	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAAG ACGAGTATTA GGAGAGCCCG TTGGTGGCTGG GCTCAGACAT	120 180 240 300 360 420 480 540 600 660
	TCCAAAGTTC CACAGAAAGTA GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGCCTCCTTC AAAACCGCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG CCTTCATCCC AATATTCCCA CCTGGCCCTC GGCGACCAAC GTGTAACATT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTTCCCCTGC CTTCCCCTGC CAATTTCTG TGCCTGCTCC TGCCTGCTAGGCAAA CTTTTAAAAT	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTCTG TCACTCCTG TCACTCCTG TCACTCTTG CCGCGCCAGC CTAGGTCTTG	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGCTGG GCTCAGACAT GTTTTGTTGA	120 180 240 300 360 420 480 540 660 720 780
50	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTTGGATGT CACAACCAAC CTCTGGCTTT TCAGGCTTGA TCTAGGCTTGA AAATGCCCTC TTTTTTCTTA	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGCCTCCTTC AAAACCGCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAAGAG	CCTAGGCATC AAGTGTACAG AGGACCGAA AGGACTATCC CATTTTTGTT GGCTGTGGAG CTCTTCATCCC AATATTCCCA CCTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCCTGC CAATTTTCTG TGCCTGCTCC ACTAGGCAAA AGAGGGACAG	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTCTG TCACTCCTGG CGCGGCCAGC CTAGGTCTTG CTAGGTCTTG CATAGAAAGT	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGCTGG GCTCAGACAT GTTTTGTTGA CCCCAAAGAG CCCCAAAGAG	120 180 240 300 360 420 480 540 600 660 720 780 840
	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTCTTA CACAAAGGTT CAGCAAAGGTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGCCTCCTTC AAAACCGCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAAGAAAT TTAAAAGAAAT	CCTAGGCATC AAGTGTACAG AGGACCGAA AGGACTATCC CATTTTTGTT GGCTGTGGAG CTGTGAGGACT CCTTCATCCC AATATTCCCA CCTGGCCCTC GGCGACCAC CTGTACATT TGATCATAAA TCACAAGCCT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTTCCCCTGC CAATTTTCTG TGCCTGCTCC ACTAGGCAAA CTTTTAAAAT AGAGGGACAG AATCTGTCAC	TCCAGTCGAG GTCAAAAAAT ACTCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CCCCAGTGCA GTCCTCTCTG TCACTCCTGG CCGGGCCAGC CTAGGCTCTTCTG CATGGCTCTTCTG TCACTCCTGT TCACTCCTGT TCACTCCTGT TCACTCCTTGT TCACTCCTTGT TCACTCCTTGT TCACTCCTTGT TCACTCTTTT	AACATGTATA GACCACAGCG CCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAGA ACGAGTATTA GGAGAGGCCG TTGGTGCTGG GCTCAGACAT GTTTTGTTGA CCCCAAAAGAG TTTGCTATTA	120 180 240 300 360 420 480 540 600 720 780 840 900
50	TCCAAAGTTC CACAGAAGTAG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTCTTA CACCAAGGTT CCAGCAAGGTT CCAGTCACAA	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT AGCCTCCTTC AAAACCGCAA TCCTCTCAGC TAGAGGCCAA TTCATTCAC AATAAAAGAC TTAAAGAAAT TTTAAACTAGG	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG CCTTCATCCC AATATTCCCA ACTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTTTG	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCCTGC CAATTTTCTG TGCCTGCTCC ACTAGGCAAA CTTTTAAAAT AGAGGGACAG AAAACTTGTT	TCCAGTCGAG GTCAAAAAAT ACCTCAGGAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG TCACTCCTGG CCGGGCCAGC CTAGGTCTTG CATAGAAAGT TGTCTTATAA TTGGTTTGCT	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGCTGG GCTCAGACAT GTTTTGTTGA CCCCAAAGAG TTTGGTCCCAA	120 180 240 300 420 480 540 600 660 720 780 840 900 960
50	TCCAAAGTTC CACAGAAGTA GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGCT TTATATCTTA CACCAAGGTT CAGCAAGGTT CCAGTCACAA GAGGCACTAG	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGCCTCCTTC AAAACCGCAA TCCTCTCAGG TAGAGGCCAA TTCATTTCAC AATAAAGAA TTTAAACTAGG CTGGGGCCCC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG CCTTCATCCC AATATTCCCA CCTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTACAGAGTGT TACAGAGTGT TACAGAGTGT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCTGC CAATTTTCTG TGCCTGCTCC TGCCTGCCAATTTTCTAAAAT AGAGGACAA ACTTTTAAAAT AGAGGACAG AATACTGTTA AGAGCAGAGC	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTG TCACTCTGG CCGCGCCAGC CTAGGTCTTG CATAGAAAGT TGTCTTATAG TTGGTTTTACTT TTGGTTTTTCTTTTC	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGCCG TTGGTGCTGG GCTCAGACAT GTTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGCAATG	120 180 240 300 360 420 480 540 600 720 780 840 900 960 1020
50	TCCAAAGTTC CACAGAAGTA GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGCT TTATATCTTA CACCAAGGTT CAGCAAGGTT CCAGTCACAA GAGGCACTAG	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT AGCCTCCTTC AAAACCGCAA TCCTCTCAGC TAGAGGCCAA TTCATTCAC AATAAAAGAC TTAAAGAAAT TTTAAACTAGG	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG CCTTCATCCC AATATTCCCA CCTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTACAGAGTGT TACAGAGTGT TACAGAGTGT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCTGC CAATTTTCTG TGCCTGCTCC TGCCTGCCAATTTTCTAAAAT AGAGGACAA ACTTTTAAAAT AGAGGACAG AATACTGTTA AGAGCAGAGC	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTG TCACTCTGG CCGCGCCAGC CTAGGTCTTG CATAGAAAGT TGTCTTATAG TTGGTTTTACTT TTGGTTTTTCTTTTC	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGCCG TTGGTGCTGG GCTCAGACAT GTTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGCAATG	120 180 240 300 420 480 540 600 660 720 780 840 900 960
50 55	TCCAAAGTTC CACAGAAAGTA GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTTCTTA CAGCAAGGTT CCAGTCACAA GAGGCACTAG TCTAGGCACAAG	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGCCTCCTTC AAAACCGCAA TCCTCTCAGG TAGAGGCCAA TTCATTTCAC AATAAAGAA TTTAAACTAGG CTGGGGCCCC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGGACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA CCTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTGTTT TACAGAGTGC GACTGAATCA	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCCTGC CAATTTCTG TGCCTGCTCA ACTAGGCAAA ACTTGTTAAAAT AGAGGGACAG AATCTGTCAC AAAACTTGTCAC AAGAATGAAC AAGAATGAAC	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTG TCACTCCTGG CGCGGCCAGC CTAGGTCTTG CATAGAAAGT TGTCTTATAA TTGGTTTGCT TTCATTTTCC CCTCTTCCCG	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGCTGG GCTCAGACAT GCTTTGTTGTA CCCCAAAGAG TTTGCTATTA TCTGCTATATA TCTGTCCCAA TCTGTCCCAA	120 180 240 300 360 420 480 540 600 720 780 840 900 960 1020
50 55	TCCAAAGTTC CAGAGAAGTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTTCTTA CAGCAAGGTT CCAGTCACAA GAGGCACTAA GAGGCACTAA TCTAGGGTCG TCTAGGGTCC TATTGGGCCCC	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT AGACCGCAA TCCTTCTCAGC TAGAGGCCAA TTCATTCAC AATAAAAGAAAT TTTAACTAGG CTGGGGCCCA AGGGACCTCA AGGCCCAAC AGCCCAGCC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGGACTATCC CATTTTTGTT GGCTGTGGAG CTCCTTCATCCC AATATTCCCA CCTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTTTG TACAGAGTGC GACTGAACAT TCTGGAAAAT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCCTTCC CTTCCCCTGC CAATTTTCTG TGCCTGCTCC ACTAGGCAAA ACTTTAAAAT AGAGGGACAG AATCTGTCAC AAAACTTGTT AGGGCAGAG AAGAATGAAG CAGCTCCTCT	TCCAGTCGAG GTCAAAAAAT ACTCTAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CCCCAGTGCA GTCCTCTCTG TCACTCCTGG CCTAGGCTCTTT CATTCTTACAT TCACTCTTT TCACTCTTTT TCACTCTTT TCACTCTTT TCACTCTTT TCACTCTTT TCACTCTTT TCACTCTTT TCACTCTTCCT CTAGGAAAGT TGTCTTATAA TTGGTTTGCT TTCACTTCTCCC CTGAGGAAAAC	AACATGTATA GACCACAGCG CCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGTA ACCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGCTGG GCTCAGACAT GTTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGATGT TGATGTTTCC AAAGTCAACT	120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020
50	TCCAAAGTTC CACAGAAGTAG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTTCTTA CACAAGGTT CCAGTCACAA GAGGCACTAG TCTAGGGTCACAA GAGGCACTAG TATGGCCCCC GAGACTGAGA	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAA TCCTCTCTGTG AGACCCCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAAGAG TTAAAGAAAT TTTAACTAGG CTGGGGCCCC AGGGACCTCA AGCCCAGCC CTGGGAGCAG	CCTAGGCATC AAGTGTACAG AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA ACTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTGTTG TACAGAGTGC GACTGAATCA AGTTGGAAAAT AGTTGGCAAA	TTATTGATCC CTGATGAGTI TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG TGCCTGCTCC ACTAGGCAAA CTTTTAAAAT AGAGGACAG AATCTGTCA AAACTTGTT AGGCAGAGC AAGAATGAAG CAGCTCCTCCTCC CTGCCAGAAG	TCCAGTCGAG GTCAAAAAAT ACCTCAGGAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGGTTGAGAC CCCCAGTGCA GTCCTCTGG TCACTCTGG CCGGGCCAGC CTAGGTCTTG CATAGAAAGT TGTCTTATAA TTGGTTTGCT TTCATTTTC CCTCTTCCCC CTGAGAAAC CCTCTCGCAT CCAGAAAC CCTCTCGCAT	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACACAGG ACGAGTATTA GGAGAGGCCG GTTGGTGCTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGGTCCCAA GTTTGAATGT TGATGTTGAATGT TGATGTTGAACACT CCTGAACACT	120 180 240 300 360 420 600 660 720 780 840 960 1020 1080 1140 1200
50 55	TCCAAAGTTC CACAGAAGTA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAT AAATGCCCTC TTTTTCTTA CAGCAAGGTT CCAGTACACA GAGGCACTAG TCTAGGCTCG TATTGGCCCCC GAGACTGAGA ATCCTGAGTA ATCCTGAGTA	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CCTGTCTGTGG AGCCTCCTTC AAAACCGCAA TCCTCTCAGCAC ATCATTCAC AATAAAGAAG TTAAAGAAG CTGGGGCCC AGGGACCTCA AGCCCCAGCC CTGGGGACCA ATTATGACCA ATTATGACCA ATTATGACCA	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA CCTGGCCCTC GGCGACCAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTTGTGTTG TACAGAGTGC GACTGAATCA TCTGGAAATCA TCTGGAAATCA TCTGGAAAAC CAAACTGCGC	TTATTGATCC CTGATGAGTGT GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTCCCTGC CAATTTTCTG TGCTGCTAAAAT AGAGGACAA CTTTTAAAAT AGAGGACAG AATCTGTCA AAAACTTGTTA AGGCAGAGC AAGAATGAAG CAGCTCCTCT CTGCCAGAGC CTGGCATTG CCTGGCATTG	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGGTTTAGAC CCCCAGTGCA GTCCTCTG TCACTCCTG CATAGAAAGT TGTCTTATAGAC CTAGGTCTTG CATAGAAAGT TTGTTTTATCT CTCTTTTCCCG CTGAGGAAAG CCTCTCGCAT GAGGAAAGC GAGGAAAGCC	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGCCG TTGGTGCTGG GCTCAGACAT GTTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCCAA GTTTGCATTT GATGTTGCT TGATGTTTGA TCTGTCCCAA GTTTGAATGT TGATGTTTTC AAAGTCAACT CCTGAACACT CACTGTGGTC CACTGTGGTC CACTGTGGTC	120 180 240 300 360 420 540 600 660 720 780 840 900 1020 1080 1140 1200 1260
50 55	TCCAAAGTTC CACAGAAAGTA GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC TTTTTTCTTA CAGCAAGGTT CCAGTCACCAA GAGGCACTAG TCTAGGCTCGC TATGGCCCCC GAGACTGAGA ATCCTGAGTA ACTGTTGAGA	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AAACCGCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAGAGA TTTAAACTAGC CTGGGGCCCC AGGGACCTCA AGCCCCAGCC CTGGGGGCCC AGCGCCCTGGAGCAC TTTAAGCAAG TTTAAGCAC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGGACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA CCTGGCCCTC GGCGACCAAC TGTAACATT TGATCATAAA TCACAAGCCT TTTTGTGTTG TACAGAGTGC GACTGAATCA TCTGGAAAAT AGTTGGCAAA AGTTGGCAAA CAAACTGCGC CAGCCTTGGT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCCTGC CAATTTTCTG TGCCTGCTCA ACTAGGCAAA ATCTGTCAC AAAACTTGTT AGGCAGAGC AAAACTGTCAC AAGAATGAAC CCTGCCAGCAG CCTGCCAGCAG CCTGCCAGCATG CCTCTCTCTA	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG CACACCCGGC CTAGGTCTTG CATAGAAAGT TGTCTTATAA TTGTTTTGCT TCATTTTCC CCTCTCCGC CTGAGGAAAC CCTCTCCCGC GAGGAAAC TCCTCTCCCG TGAGGAAAC TCCTTCCCGC TGAGGAAAC TCCTTCCGCAGAGAAC CCTCTCCGCAG	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGCTGG GCTCAGACAT CCCCAAAGAG TTTGCTATTA TCTGCTATTA TCTGTCCCAA GTTTGCTATTA TCTGTCCCAA GTTTGAATGT TGATGTTGTC AAAGACT CCCGAACACT CACTGTGGTC CACTGTGGTC GGAATACACC	120 180 240 300 360 420 480 600 660 720 840 900 900 1020 1080 1140 1200
50 55	TCCAAAGTTC CACAGAAGTAG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATTG GCTGCTCTT TCAGGCTGAC AAATGCCCTC TTTTTTCTTA CACAAGGTT CCAGTCACAA GAGGCACTAG TCTAGGGTCG TATGGCCCC GAGACTGAGA ACTGTTGAGA ACTGTTGAGA	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGCCTCCTTC AAAACGCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAAGAT TTTAACTAGG CTGGGGCCCC AGCGACCTCA AGCCCCAGCC CTGGGAGCAG ATTATGACCAA TCTCCGTCAA	CCTAGGCATC AAGTGTACAG AGCACTATCC CATTTTTGTT GGCTGTGGAG CCTTCATCCC AATATTCCCA ACTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAACAT TCACAAGACT TTTTGTTT TACAGAGTGC GACTGAAACT AGTTGGCAAA AGTTGGCAAA CGACTGAATCA CAAACTGGCC CAGCCTTGGT	TTATTGATCC CTGATGAGTI TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CATTTCTG TGCCTGCTCAAAAACTTGTTAAAAT AGAGGGACAA AAACTTGTTA AGAATGAAG CAGATCCTCTC CTGCCAGAAG CTGCCAGAAG CTGCCAGAAG CAGCTCCTCT CTGCCAGAAG CCTTGCCATTA GACGAACGCC	TCCAGTCGAG GTCAAAAAAT ACTCCAGTAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG CCGGCCAGC CTAGGTCTTAGAAAGT TGTCTTATAAA TTGGTTTGCT TCCATTTTCC CCTCTTCCCG CTGAGGAAAC CCTCTCGCAT GAGAGAAAGC CTCTCGCAT TCAGACAT TCTGTTACAA	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGCTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGATGTT TGATGTTGTC AAAGTCAACT CCTGAACACT CCTGAACACT CCTGAACACT CACTGTGGTC CGGAATACACC CGACACCCTTT	120 180 240 360 420 480 540 660 720 780 840 900 960 1020 1140 1260 1260 1320
50 55 60	TCCAAAGTTC CACAGAAGTAC CACTGGCTGC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTTTTTATA CAGCAAGGTT CCAGTCACAA GAGGCACTAG TCTAGGCTCA TATGGCCCCC GAGAACTGAGA ACTGTTGAGA ATTGCACTC	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAA TCCTCTCTGTGG AGCCTCCTTC AAAACGCAA TCCTCTCAGC TTAGAGGCCAA TTCATTTCAC AATAAAAGAT TTTAACTAGG CTGGGGCCCC AGGGACCTCA AGCCCCAGCC CTGGGAGCAG ATTATGACCA TTCTCTCCGTCAA TCTTCTCCCCCA TTCTGAATGG	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTGTT GGCTGTGGAG CCTTCATCCC AATATTCCCA ACTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTTTG TACAGAGTGC GACTGAATCA ACTGGAAAAAT AGTTGGCAAA CAAACTGCGC CAGCCTTGGT CAATGTGTTG CAATGTGT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGC CCTTCCCCTGC CAATTTTCTG TGCCTGCTCC ACTAGGCAAA CTTTTAAAAT AGAGGGACAG AATCTGTCA AAAACTTGTT AGGCAGAGC CAGCAGAAG CCTGCCTCT CTGCCAGAAG CCTGCCTCTCT CTGCCAGAAG CCTGCATTG CCTCTCTCT AGCGAACGCC AGCAAGCCC AGCCAGCTAT	TCCAGTCGAG GTCAAAAAAT ACTCAGAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG TCACTCTGG CCGGGCCAGC CTAGGTCTTG CATAGAAAGT TTGTTTATTC CCTCTTCCCG CTGAGGAAAC CCTCTCGCAT GAGAAAGC CTCCTGAGAAA GAGAGCCGGA	AACATGTATA GACCACAGC CCCCCTTTGT GAGTTTCATA ATCTTCTTGT GGCAGCCAGT ACCACACAAG ACGAGTATTA GGAGAGGCCG GTTGGTGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGATGT TGATGTTGTC CAAGTCAACT CACTGTGGTC GGAATACACT CACTGTGGTC CGACACCTTT CACCTTTTTT	120 180 240 300 360 420 600 660 720 780 840 900 1020 1140 1200 1260 1320 1380 1440
50 55	TCCAAAGTTC CACAGAAGTAC CACTGGCTGC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTTTTTATA CAGCAAGGTT CCAGTCACAA GAGGCACTAG TCTAGGCTCA TATGGCCCCC GAGAACTGAGA ACTGTTGAGA ATTGCACTC	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGCCTCCTTC AAAACGCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAAGAT TTTAACTAGG CTGGGGCCCC AGCGACCTCA AGCCCCAGCC CTGGGAGCAG ATTATGACCAA TCTCCGTCAA	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTGTT GGCTGTGGAG CCTTCATCCC AATATTCCCA ACTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTTTG TACAGAGTGC GACTGAATCA ACTGGAAAAAT AGTTGGCAAA CAAACTGCGC CAGCCTTGGT CAATGTGTTG CAATGTGT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGC CCTTCCCCTGC CAATTTTCTG TGCCTGCTCC ACTAGGCAAA CTTTTAAAAT AGAGGGACAG AATCTGTCA AAAACTTGTT AGGCAGAGC CAGCAGAAG CCTGCCTCT CTGCCAGAAG CCTGCCTCTCT CTGCCAGAAG CCTGCATTG CCTCTCTCT AGCGAACGCC AGCAAGCCC AGCCAGCTAT	TCCAGTCGAG GTCAAAAAAT ACTCAGAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG TCACTCTGG CCGGGCCAGC CTAGGTCTTG CATAGAAAGT TTGTTTATTC CCTCTTCCCG CTGAGGAAAC CCTCTCGCAT GAGAAAGC CTCCTGAGAAA GAGAGCCGGA	AACATGTATA GACCACAGC CCCCCTTTGT GAGTTTCATA ATCTTCTTGT GGCAGCCAGT ACCACACAAG ACGAGTATTA GGAGAGGCCG GTTGGTGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGATGT TGATGTTGTC CAAGTCAACT CACTGTGGTC GGAATACACT CACTGTGGTC CGACACCTTT CACCTTTTTT	120 180 240 360 420 480 540 660 720 780 840 900 960 1020 1140 1260 1260 1320
50 55 60	TCCAAAGTTC CACAGAAGTAC CACTAGACAAC CTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAT TCAGGCTGAT CACAAGGTT CCAGTACACA GAGGCACTAG TCTAGGCTCG TATTGGCCCC GAGACTGAGA ATCCTGAGTA ACTGTGACAAC ACTGTGACAA ACTGTGACAT ACTGTTGACATC AGAGTCTTTGACATCA AGAGTCTTTGACATCA AGAGTCTTTGACATCTA AGGAATTCTTA AGGAATTCTA	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT AGACCCCAA TCCTCTCAGC AGACCCCAA TTCAATTCAC AATAAAGAAG TTTAAACTAGG CTGGGCCCC AGGGACCTCA AGCCCCAGCC ATTATGACTAA ATTATACTAGG ATTATGACTAA AGCCCCAGCC AGCGACCTCA AGCTCCACTCAA TCTTCTCCCAA TCTTCTCTCCAA AGAGGACCCA AGAGGACCCAA	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACCCTC AATATCCCA ATATCCCA ATATCCCA CTGGCCCTC GTGTAACAT TGATCATAAA TCACAAGCCT TTTTGTGTTG TACAGAGTGC GACTGAATCA ACTGGAATCA CTGGAATCA ACTGGAATCA CAATGCGC CAGCCTTGGT CACCTGGTA CAATGTGGC CAGCCTTGGT CAATGTGGC CAACTGGTG CAATGTGGC CAACTGGTG CAATGTGGC CAACTGGTG CAATGTGGTG CAATGTGGTG CAAGCATGAG	TTATTGATCC CTGATGAGTGT GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG TGCTGCTAA ATTTAAAAT AGAGGACAG AATCTGTCA AAAACTTGTTA AGAGCAGAGC	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTCGTTTG GTCATCTGGT TCACTCTGGT CACTCCTGG CTAGGTCTTG CATAGAAAGT TGTCTTATTC CTATAGAC TTCATTTTC CTGTTTCCCG CTGAGGAAAG CCTCTCGCAT CAGGAAAGC TCCTTAGACAT CAGGAAAGC TCCTAGACAT CTGTTACAG GAGAAAGC CCACCAGAC CCAACCAGAT	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGCCG TTGGTGCTGG GCTCAGACAT GTTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGCATTT GAAGTCAACT CCTGAACACT CACTGTGGTC GGAATACACC CGACACCTTTT CACCTTTTT GGTCCCGATC	120 180 240 300 360 420 600 660 720 780 840 900 1020 1140 1200 1260 1320 1380 1440
50 55 60	TCCAAAGTTC CACAGAAAGTA GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC TTTTTTCTTA CAGCAAGGTT CCAGTCACCAA GAGGCACTAG TCTAGGCTCGC TATGGCCCC GAGAACTGAGA ATCCTGAGTA ATCTGAGAA ATCTGAGAA ATTGACATCA GAGTCTCTTG GAGAATTCTA AGGAATTCTA ATCAAGGATG	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AAACCGCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAGAG TTTAAACTAGC CTGGGGCCCC AGGGACCTCA AGCCCCAGCC CTGGGAGCAG TTCTCTCACA TCTCTCCCA TCTCTCACA TCTCTCACA CTTCTCCCA TCTCTGAATGG GAGAGCCCA GCAAGGGCCCA GCAAGGGCCCA GCAAGGGCCCA CCCGCACCC CTGGAACCAG CCTCGAACA CCTCTCAATGG CCAAGGACCCA GCAAGGGCCCA GCAAGGGCCCA GCAAGGGCCCA	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA CCTGGCCCTC GCCGACCAAC TGTAACATT TGATCATAAA TCACAAGCCT TTTTGTGTTG GACTGAATCA TCTGGAAAAT AGTTGGCAA AGTTGGCAA ACAACTGCGC CAGCCTTGGT GACCTGGTAC CAATGTGGTG CCAGCATGAGG GCATGAATAG GTACACATT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CATTTCTG TGCCTGCTCA ACTAGGCAAA ATCTGTCAC AAAACTTGTT AGGCAGAGC AAAACTGTCT CTCCCAGAGC AGAATGAAC CCTGCCAGAGC AGCCAGCTAT CCTCTCTCTA AGCGAACGC AGCCAGCTAT ATCACCATGC AGGAACCC AGCCAGCTAT ATCACCATGC AGGAATGACC AGCAAGCC AGCAAGCACA	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG TCACTCCTGG CATAGAAAGT TGTCTTATAA TTGGTTTGCT TTCATTTTCC CTGAGGAAAC CCTCTCGCAG CCTCTCCGCAG CCTCTCCGCAG TCCTTCCCG TGAGGAAAC TCCTTAGACAT TCGTTACAA GGATCCCGGA TCCTAGACAT TTGTTACAA CCAACCAGAT TTGATGCCGG	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGCTGG GCTCAGACAT CCCCAAAGAG TTTGCTATTA TCTGCTATTA TCTGTCCCAA GTTTGCTATTA TCATGTCCCAA GTTTGAATGT TGATGTTGTC CAACACT CACTGTGGTC GGAACACT CACTGTGTC GGAACACT CACCTTTTT CACCTTTTT CACCTTTTT CACCTTTTT CACCTTTTT CACCTTCCCCAA AGGCCCTTT CACCTTCCCCAACACT CACCCTTTT CACCTTCTTT CACCTTCTTT CACCTTCTTT CACCTTCTCT ATGCTCACTC	120 180 240 300 360 420 540 600 660 720 780 840 900 1020 1140 1260 1320 1320 1340 1500
50 55 60	TCCAAAGTTC CACAGAAGTTG CACAGAAGTAG CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTTTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAA TCCTCTCTGC AGACCCAA TCCTCTCAGC TAGAGGCCAA TTCATTCAC AATAAAAGAC TTAAAGAAAT TTTAACTAGG CTGGGGCCCC AGGGACCTCC CTGGGAGCAG ATTATGACCA TCTCCGTCAA TCTTCTCCCCA TCTCGTCAA TCTTCTCCCCA TCTGAATGG AGAGGACCCCA GCAAGGGGTTT GAATTCCAAT GCAAGGTGTT GAATTCCAAT	CCTAGGCATC AAGTGTACAG AGCACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG CCTTCATCCC AATATTCCCA ACTGGCCCTC GGCGACCAAC GTGTAACATT TGATCAAGCCT TTTTGTGTT TACAGAGTG GACTGAATA AGTTGGCAAA CCAGCCTTGGTCCAAACTGGCCAC CAGCCTTGGTCCAAACTGGCCAAC CAAACTGGCCAAC CAAACTGGCCAAC CAACTGGGTAC CAATGTGGTG CAATGTGGTG CCAGCATGAGT CCAACATGAGTG CCAGCATGAGT CCAACATGAGTG CCAGCATGAGT CGACATGAGT GGTACACAAT GGATCTCAC	TTATTGATCC CTGATGAGTI TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG ACTGGCAAA CTTTTAAAAT AGAGGGACAA AAACTTGTT AGGCAGAGC AAGAATGACC CTGCCAGAAG CCTGCCTCTCTCTC CTGCCAGAAG CCTGCCATTCACAGCAC ACGAGCACTAT ATCACCATGC AGGATGACCA TCTTCCCTCT CTGCCATGC AGGATGACCA CTTTCACCATGC AGGATGACCA TCTTGCCCTC	TCCAGTCGAG GTCAAAAAAT ACCTCAGTAG CACCATCGGT ATTCTTACTA ACTCCTGTTA CTCCTGTTG CCCCAGTGCA GTCCTCTGG TCACTCCTGG CCGGGCCAGC CTAGGTCTT TTCATTTTC CCTCTTATAA TTGGTTTGCT TTCATTTTC CCTCTTCCCGA CCAGGAACACAGAT TCGGTCACAA GGATCCCGA CCAACCAGAT TTGATCCCGT TTGATCCCGT TCAGTCCCGA TCTGACACT TCTGTCCCGA TCTGACACT TCTGTCCCGA TCTGACCACT TCTGACCACT TCTGACCCGA TCTGACCACT TTGATCCCGG TACCTTTCCC	AACATGTATA GACCACAGG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGTGGG GCTCAGACAT GTTTGTTGA CCCCAAAGGA GTTTGAATGT TGATGTGTCCAA GTTTGAATGT TGATGTTGACCC CAAGACACT CCTGAACACT CCTGAACACT CCTGAACACT CACTGTGGTC CGAACACT CACTGTGGTC CGACACCTTT CACCTTTTTT GGTCCGCATC ATGCTCACCT TAGCTTTTCC TAGCTTTTCC	120 180 240 300 360 420 600 720 780 840 900 960 1020 1140 1260 1320 1380 1440 1500 1560
50 55 60	TCCAAAGTTC CACAGAAGTAC CACTAGACAAC CTTTGCCTCT CCTTGGCAGA TTTTCTTGGC CTCTGGCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTTTTTA CAGCAAGGTT CCAGTCACAA GAGGCACTAG ATCTGGCTCA AATGCCCCC GAGACTGAGA ATCTGAGTA ACTGTTGAGA ATTGACATCA AGGTATCATTGACAAC AATGACTCTTTG AGGAATTCTA GAGTATCTTTG AGGAATTCTA TACAAGGATG CACATGGTAGA ATTGACATCA ATTGACATCA AGTTCTTTTTTTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT AGCCTCCTTC AAAACCGCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAAGAA TTTAACTAGG CTGGGGCCCC AGGGACCTCA AGCCCCAGCC CTGGGAGCAG ATTATGACCA TTTTCTCCCCA TTCTCTCCCCA TTCTTCTCCCCA TCTCGAATGG AGAGGACCCA GCAAGGTGTT AATGAAGATA ATTACAAT	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTGTT GGCTGTGGAC CCTCATCCC AATATCCCA ACTATCCC AATATCCCA CCTGGCCCTC GGCGACCAAC GTGTAACAT TGATCATAAA TCACAAGCCT TTTTGTGTTG TACAGAGTGC GACTGAAAA CAAACTGCGC CAGCCTTGGT CCAATGTGGC CAATGTGGC CAATGTGGC CAATGTGGT CCAATGTGGC CAATGTGGT CCAATGTGGT CCAATGTGGT CCAATGTGGT CCAATGTGGT CCAATGTGGT CCAATGTGGT CCAACTCAC CTACAAGTGG CTACAAGTGG	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGC CCTGCCTTCC CATTTCTG TGCCTGCTCC ACTAGGCAAA CTTTTAAAAT AGAGGGACAG AATCTGTCA AAAACTTGTT AGGCAGAGC CAGCTCTCT CTGCCAGAAG CCTGCCTCTCT CTGCCAGAAG CCTGGCATTG CCTCTCTCTA AGCAACGCC AGCAAGCTCATC ATCACCATGC AGCAAGCTC TTGCCCTC TGAAAATTTCA	TCCAGTCGAG GTCAAAAAAT ACTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTCTCTGGT CACTCCTGGT CACTCCTGGT CACTCCTGG CCGGGCCAGC CTAGGTCTTG CATAGAAAGT TTGGTTTACC CTCTGTCCCG CTGAGGAAAC CCTCTCGCAT GAGAAAGC TCCTAGAAA GATCCCGGA CCAACCAGAT TTGATCCCGA CCAACCAGAT TTGATCCCGA CAACCAGAT TTGATCCCGA TCTGATCCCGA CCAACCAGAT TTGATCCCGA CAACCAGAT TTGATCCCGA TTGATCCCGA CAACCAGAT TTGATCCCGA TTGATCCCGA CAACCAGAT TTGATCCCGA CAACCAGAT TTGATCCCGA CACTCTTCTCC AGCTTGAAAT	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTG GGAGCCAGT ACCACACAAG ACGAGTATTA GGAGAGGCCG TTGGTGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGATGT TGATGTTGTC CAAGTCAACT CACTGTGGTC GGAATACACT CACTGTGGTC CGACACCTTT CACCTTTTT GGTCCGCATC ATGCTCATC ATGCTCATC TAGCTTTTC CACTTTTTT CTGCCCATC CACACCTTT CACCTTTTT CTGCCCATC CACACCTTT CACCTTTTTT CTGCCCCATC CACACCTTT CACCTTTTTC CTAGCTTTTCC CAATGAGAAG	120 180 240 300 360 420 6600 6600 720 780 840 900 1020 1140 1200 1260 1380 1440 1560 1560 1620 1680
50 55 60 65	TCCAAAGTTC CACAGAAGTAG GTGTAAAGAG CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGCT TTTTTCTTA CACCAAGGTT CCAGTCACCAA GAGGCACTAG TCTAGGCTCC GAGACTGAGA ATCCCTGAGTA ACTGTTGACATCA AGTGACATCA CACAGGTTT ACAAGGAT CACAGGTT ACTGTTGACA ATTGACATCA AGTGACTCAT TACAAGGATG CACATGCTCA TACAAGGATG TACAAGGATG ATTCCTGAGA ATTCCTGAGA ATTCCTGAGA	TTCCAGTCCT CTCAAATCAT AGCCAAATCA CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGACCCCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAGAG TTTAAACTAGG CTGGGGCCCC AGGGACCTCA AGCCCCAGCC ATTATGACCA TTTTCCCAA TCTTCTCCCA TCTTCTCCCA TCTTCAGTCAA TCTTCTCCCA TCTTGAATGG CTAGAGGCCCA TCTTCAACT TCTGAATGG ATTATGACCA TCTTCACAT TCTGAATGG AGAGGACCCA TCTTGAATGG AGAGGACCCA TCTGAATGG AGAGGACCCA TCTGAATGG AGAGGACCCA TCTGAATGG AGAGGACCCA TCTGAATGAT AGCACTTCCAA	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACCCTC CATATCCCA AATATCCCA ACTATCCC AATATCCCA CCTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TATCGGAATCA TCTGGAATCA TCTGGAATCA TCTGGAAAAA CAAACTGCGC CAGCCTTGGT GACCTGGTAC CAACTGGTAC CCAGCATGAG GTACACAATT GGATCTCAC GTACACAATT GCATCCACACTCC CTACACAGTGC CTACACAGTGC GTACACAATT TCACAAGTGC CTACACAGTGC CTTTGATTTT	TTATTGATCC CTGATGAGTGT GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTTCCCTGC CAATTTTCTG TGCTGCTCC CACTAGGCAAA CTTTTAAAAT AGAGGGACAG AATACTGTTA AGAGCAGAGC	TCCAGTCGAG GTCAAAAAAT ACCTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTG TCACTCTGG TCACTCTGG CTAGGTCTTG CATAGAAAGT TGTCTTATAC TTCATTTTC CCTCTTCCG CTGAGGAAA CCTCTGGCAT TCGTTACAA GGATCCCGA CCACCAGAT TCGTTACAA TCGTTACAA TCGTTACAA TCGTTACAA TCGTTACAA TCGTTACAA TCGTTACAA TCGTTACAA TCGTTACAA TCGATCCGG TATCTTTCT TCACTTTCCAA TCGATCCGGA TATCTTTCT TCACTTTCACAA TCGATCAGAAAAAC GCAACAAAAAC	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGCCG TTGGTGCTGG GCTCAGACAT GTTTGTTGTTGA CCCCAAAGAG TTTGCTATTA CTGTCCCAA GTTTGCAATG TCTGTCCCAA GTTTGAATGT TGATGTTGC CAAGACACT CACTGTGGTC GGAACACTT CACCTTTTT GGTCCGCATC ATGCTCACT TAGCTTTTC ATGCTTCC CAATGAGAAG TGAAATAATC	120 180 240 300 360 420 540 600 660 720 780 840 900 1020 1140 1260 1320 1320 1340 1560 1560 1680 1740
50 55 60	TCCAAAGTTC CACAGAAGTTG CACAGAAGTAG CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTTTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCTTTAA TGTGGTGCAT TCTCTCTGTGG AGCCTCCTTC AAAACGCAA TCCTCTCAGC TTGAGGCCAA TTCATTCAC AATAAAAGA TTTAACTAGG CTGGGGCCCC CTGGGAGCAC ATTATATACACA ACCCCAGCC CTGGGAGCAG TCTCTCCCA TCTCCCA TCTCTCCCA TCTCAGCTCAA GAGGACCTC AGAGGACCTCAAACACAC TCTCAGATGAA TCTCAGATGAA TCTCAGATGAA TCTCAGATGAA TCTCAGATGAA TCTCAGATGAA TCTCAGATGAA TCTCAGATGAA TCTCAGATGAA TCTCAGATGAT TTGAGATGAT TTGAGATGACTTT	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GCTGTGGGGGGCTC CCTTCATCCC AATATTCCCA AGGACCAAC GTGTAACATT TGATCATACCT TTTGTTTG TCACAAGCCT TTTTGTTTG TACAGAGTGC GACTGAAAT AGTTGGCAAA CGAGCATGAA CCAGCCTTGGT CAACTGGGC CAGCCTTGGT GACCTGGTAC CAACTGGTCA CTACAAGTGG GTACACAATT GGATTCTCAC CTACAAGTGG GTTCACATTT CCTACAAGTGG CTACAAGTGG CTACAAGTGG CTACAAGTGG CTACAAGTGG CTACAAGTGG CTACAAGTGG CTTTGATTTT CATGGTCATG	TTATTGATCC CTGATGAGTI TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC ACTAGGCAAA CTTTTAAAAT AGAGGACAG AAACTTGTCA AAAACTTGTTA AGAGAAGAG CCTGCCATCC CTGCCATGC CAGCAAG CCTCTCTCT CTGCCAGAAG CCTCTCTCTAGAACC ACAGAACCC ACCAGCATAT ATCACCATG ACGAATTTCC ACAGAGGAGAC AGGATGACCA ACAGGAGTGA ACAGAGTGA ACAGGAGTGA ACAGAGTGA ACAGGAGTGA ACAGATTTCT	TCCAGTCGAG GTCAAAAAAT ACCTCAGAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGTG CCCAGTCCTGC CTAGGTCTTG CATAGAAAGT TTGTTTATAA TTGGTTTGCT TTCATTTTTC CCTCTTCCGC CTGAGGAAAC CCTCTCGCAT GAGAGAAAC CCTCTCGCAT TCTGTTACAA GGATCCCGGA TCTGTTACAA GATCCCGGA TTGTTACAA GATCCCGGA TTGTTACCAG TTGATGCCGG TATCTTTCTC AGCTTGAAAT GCAACAAAAA CCAAT CCAACAAAAA CCAAT TCAATGTGAA TCAATGTGAA	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACACAAG ACGAGTATTA GGAGAGGCCG TTGGTGGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGATGTTC CAAGTGTTGTC CAAGACACT CACTGTGGTC CGGAACACT CACTGTGGTC CGGAACACT CACTGTGTTC CACGCGTTT CACGCGGTTT CACGCGGTTT	120 180 240 300 360 420 480 600 660 720 840 900 900 1020 1140 1220 1380 1440 1500 1560 1620 1
50 55 60 65	TCCAAAGTTC CACAGAAGTTC CACAGAAGTAG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC GCTCTCTTT CACACAACCAC CTCTGGCTTT TCAGGCTGAT AAATGCCCTC TTTTTTCTTA CACAAGGTT CCAGTCACAA GAGGCACTAG TCTAGGGTCACAA ACTCTGAGTA ACTGTTGAGA ATTGCCTCT AGGAATTCTTA AGGAATTCTTA AGGAATTCTTA AGGAATTCTTA AGGAATTCTTA AGGAATTCTTA AGGAATTCTTA ACAAGGAT TCACAGGAT ACTCTGAGA AATTCCTGAGA AACTCCTGAGA AACTCCTGAGA AACTCCTGAGA AACTCCTGAGA AACACCCCAG	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAA TCTCTCTGTGG AGCCTCCTTC AAAACGCAA TCCTCTCAGC TTAGAGGCCAA TTCATTTCAC AATAAAAAAT TTTAACTAGG CTGGGGCCCC AGGGACCTCA ACCCCAGCC CTGGGAGCAG ATTATGACCAA TCTTCTCCCCA TCTCGTCAA TCTTCACCA TCTGAATGG AGAAGGTGTT AGCACAGTC TAGAGGACCCA TCTGGAAGGACCCA TCTGGAATGG AGAAGGTGTT AGCACAAT ATGAGATGAT ATGAGATGAT ATGAGATGAT TTGGTGACTT CCCTTCAAAA	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA ACTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTGTTG TACAGAGTG GACTGAATCA AGTTGGCAAA CGAGCATTGGC CAGCCTTGGT GACCTTGGT CAATGTGGT CCAGCATGAG GTACACATT CCTACAAGTGG CTACTGTTCAC CTACAGTGG CTACGTCAC CTTACAGTGG CTTACACTCCC CTACAAGTGG CTTACACATT CCTACAAGTGG CTTACACATC CTACAAGTGG CTTACACATC CTACAAGTGG CTTACACC CTACAAGTGG CTTACACC CTACAAGTGG CTTACACC CTTACATCC CTTACATC CTTACATCC CTTACATC CTTACATCC CTTACATCC CTTACATCC CTTACATCC CTTACATCC CTTACATCC CTTA	TTATTGATCC CTGATGAGTI GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG TGCCTGCTCC ACTAGGCAAA CTTTTAAAAT AGAGGACAG AATCTGTCA AAACTTGTT AGGCAGAGC CAGCAGAAG CCTGCCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTC	TCCAGTCGAG GTCAAAAAAT ACTCCAGTAG ACACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG TCACTCCTGG CCGGGCCAGC CTAGGTCTTG TTGTTTATAA TTGGTTTGCT TTCATTTTC CCTCTTCCGCAT GAGAGAACC CCTCTAGACAT GGATCCCGA GCACCAGAT TTGATCCCGA TCTGTTACCA TCTGTCCCGA TCTTGCCAT TCGGTTACAA GGATCCCGA CCAACCAGAT TTGATGCCGG TATCTTTCTC AGCTTGAAAAC CCAACAAAAAC CCAACAAAAAC CCACGATGCT TCAATGTGAG CCACGATGCT TCAATGTGAG CCACGATGCT	AACATGTATA GACCACAGG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACACAGG ACGAGTATTA GGAGAGGCCG TTGGTGTGGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGGTATTA TCTGTCCCAA GTTTGATGT CACTGTGGTC CAAGACACT CACTGTGGTC CGAAACACT CACTGTGGTC CGACACCTTT CACCTTTTT GGTCCGCATC ATGCTCACCT TAGCTTTTCC CAATGAGAAG TGAAATAATC CCAGGCGTTT CCACGGGTTT CTCCTGGGTT CTCCTGGGTT	120 180 240 300 360 420 600 660 720 780 840 960 1020 1140 1200 1320 1320 1380 1440 1560 1560 1680 1740 1800 1800
50 55 60 65	TCCAAAGTTC CACAGAAGTTC CACAGAAGTAG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC GCTCTCTTT CACACAACCAC CTCTGGCTTT TCAGGCTGAT AAATGCCCTC TTTTTTCTTA CACAAGGTT CCAGTCACAA GAGGCACTAG TCTAGGGTCACAA ACTCTGAGTA ACTGTTGAGA ATTGCCTCT AGGAATTCTTA AGGAATTCTTA AGGAATTCTTA AGGAATTCTTA AGGAATTCTTA AGGAATTCTTA AGGAATTCTTA ACAAGGAT TCACAGGAT ACTCTGAGA AATTCCTGAGA AACTCCTGAGA AACTCCTGAGA AACTCCTGAGA AACTCCTGAGA AACACCCCAG	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAA TCTCTCTGTGG AGCCTCCTTC AAAACGCAA TCCTCTCAGC TTAGAGGCCAA TTCATTTCAC AATAAAAAAT TTTAACTAGG CTGGGGCCCC AGGGACCTCA ACCCCAGCC CTGGGAGCAG ATTATGACCAA TCTTCTCCCCA TCTCGTCAA TCTTCACCA TCTGAATGG AGAAGGTGTT AGCACAGTC TAGAGGACCCA TCTGGAAGGACCCA TCTGGAATGG AGAAGGTGTT AGCACAAT ATGAGATGAT ATGAGATGAT ATGAGATGAT TTGGTGACTT CCCTTCAAAA	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA ACTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTGTTG TACAGAGTG GACTGAATCA AGTTGGCAAA CGAGCATTGGC CAGCCTTGGT GACCTTGGT CAATGTGGT CCAGCATGAG GTACACATT CCTACAAGTGG CTACTGTTCAC CTACAGTGG CTACGTCAC CTTACAGTGG CTTACACTCCC CTACAAGTGG CTTACACATT CCTACAAGTGG CTTACACATC CTACAAGTGG CTTACACATC CTACAAGTGG CTTACACC CTACAAGTGG CTTACACC CTACAAGTGG CTTACACC CTTACATCC CTTACATC CTTACATCC CTTACATC CTTACATCC CTTACATCC CTTACATCC CTTACATCC CTTACATCC CTTACATCC CTTA	TTATTGATCC CTGATGAGTI GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG TGCCTGCTCC ACTAGGCAAA CTTTTAAAAT AGAGGACAG AATCTGTCA AAACTTGTT AGGCAGAGC CAGCAGAAG CCTGCCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTCTC	TCCAGTCGAG GTCAAAAAAT ACTCCAGTAG ACACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG TCACTCCTGG CCGGGCCAGC CTAGGTCTTG TTGTTTATAA TTGGTTTGCT TTCATTTTC CCTCTTCCGCAT GAGAGAACC CCTCTAGACAT GGATCCCGA GCACCAGAT TTGATCCCGA TCTGTTACCA TCTGTCCCGA TCTTGCCAT TCGGTTACAA GGATCCCGA CCAACCAGAT TTGATGCCGG TATCTTTCTC AGCTTGAAAAC CCAACAAAAAC CCAACAAAAAC CCACGATGCT TCAATGTGAG CCACGATGCT TCAATGTGAG CCACGATGCT	AACATGTATA GACCACAGG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACACAGG ACGAGTATTA GGAGAGGCCG TTGGTGTGGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGGTATTA TCTGTCCCAA GTTTGATGT CACTGTGGTC CAAGACACT CACTGTGGTC CGAAACACT CACTGTGGTC CGACACCTTT CACCTTTTT GGTCCGCATC ATGCTCACCT TAGCTTTTCC CAATGAGAAG TGAAATAATC CCAGGCGTTT CCACGGGTTT CTCCTGGGTT CTCCTGGGTT	120 180 240 300 360 420 480 600 660 720 840 900 900 1020 1140 1220 1380 1440 1500 1560 1620 1
50 55 60 65	TCCAAAGTTC CACAGAAGTAC CACTAGAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGA AAATGCCCTC TTTTTTCTTA CAGCAAGGTT CCAGTCACAA TCTAGGCTCG TATTGGCCCCC AATGCCCTCAAA ACTGTTAGAACAAA ACTGTTAGAATA ACTGTTTAGAATA ACTGTTTAGAATCA AGAGTCTCTTT AGGAATTCTA TACAAGGATG CACATGCTCA AATGCCCCAGA AACCCCAGG AACTCCTGGAAAACCCCAGG ACAACCCCAGG ACATGCTTG TCCTTTTTGGA	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT AGCCTCCTTC AAAACCGCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAGAG TTAAAGAAG CTGGGGCCCC AGGGACCTCA AGCCCCAGCC CTGGAGCAGC ATTCTCTCCCA ATTCTCCCCA TTCTCTCCCA ATTCTCCCCA TTCTGAATGG AGAGGACCCA GCAAGGTGTT GATTTCCAT ATCAGATGAT AGCTCTTCCA TTCGGTAAT CTTTCTCCAT ATCAGATGAT ATCAGATGAT ACCTTTCCAAT ACCAGACAGAA	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTGTT GGCTGTGGAG CCTCATCCC AATATCCCA ATATCCCA ATATCCCA CCTGGCCCTI GGCGACAC GTGTAACAT TGATCATAAA TCACAAGCCT TTTTGTGTTG TACAGAGTGC GACTGAAAA ACTGGAAAA CAAACTGCGC CAGCCTTGGT CCAATGTGGT CCAATGTGGT CCAATGTGGT CCTACAAGTGG GTTCACAATT GGATTCTCC CTTCAAGTGG GTTTGATTT CATGGTCATT CATGGTCATC CTTCAAGTGG CTTAGATTT CATGGTCATC CTTACAAGTGG CTTTGATTTT CATGGTCACC CTTCAATTCCCC CTTCACTCCC	TTATTGATCC CTGATGAGTTT TGTGAGCAGG GGACTTCTAA TAATTCAAGTTG CCTGCCTTGC CAATTTTCTG TGCCTGCT CACTAGGCAAA CTTTTAAAAT AGAGGGACAG AAACTTGTTA AGAGCAGAGC	TCCAGTCGAG GTCAAAAAAT ACTCAGAAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTCTCTGTG CCCCAGTGCA GTCCTCTGG CCGGGCCAGC CTAGGTCTTG CATAGAAAGT TTGTTTATC CTCTTCTCG CTGAGCAAG CCTCTCGCAT GAGAAAGC TCCTAGACAT CTGTTACAG GAGACAGAT TTGATCCGG TCCTGGCAT TCCTGTTACAG CCAACCAGAT TTGATCCCGG TAGTTCCCGAT TCTGTTACAG CCAACCAGAT TTGATCCTGA CCAACCAGAT TTGATCTCT TCTCTTCCCAC TCCAACCAGAT TTGATCTCT TCCAACTCTGAAAT CCAACAAAAAC CCAACGATGCT CCTCTAGGAT CCACGATGCT CCTCTAGGAT CCACGATGCT CCTCTAGGAT CCACGATGCT CCACGATGCT CCTCTAGGGAT CCACGATGCT CCTCTAGGGAT	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTG GGAGCAGT ACCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGATGT TGATGTTGT CAAGTCAACT CACGACACT CACGACACT CACCTTTT GGTCCGCAT ATGCTCATC CACCTTTTT GGTCCGCATC CACCTTTTT CACCTTTTTC CACGTTTTCC CAATGACACT CACGGGGTT CACGGGGTT CACGGGGTT CCTCGGGTT CACCTCTGTT CACCTCTGTT CACCTCTGTT CACCTCTTT	120 180 240 300 360 420 600 660 720 780 840 960 1020 1140 1200 1320 1320 1380 1440 1560 1560 1680 1740 1800 1800
50 55 60 65	TCCAAAGTTC CACAGAAAGTA GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAT TCAGGCTGAT TCAGGCTGAT CCAGCAAGGTT CCAGCAAGGTT CCAGCAAGGTT CCAGTCACCAA GAGCACTAG TCTAGGCTCC GAGACTGAGA ATCCTGAGTA ACTGTTGACAT ACTGTTGACAT TACAAGGATG CACATGCTCA TACAAGGATG ACTCCTGGA AACTCCTGGA AACTCCTGGA AACTCCTGGA AACTCCTGGA AACTCCTGGA ACTCCTGGA ACTCCTGGA ACTCCTGGA ACTCCTGGA ACTCCTTGGA CCATGCTCA TCCTTGTGAC TCTTTTTGG TCCTTTTGGAC TCTTTTTGGACACATGAC CCACCATGAC	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC AGCCAAATCAC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGACCCCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAGAAT TTTAAACTAGG CTGGGGCCCC AGGCCCCAGCCC AGCCCCAGCCC ATTCTTCTCCCAA TTCTTCTCCCAA TTCTTCTCCCA TTCTGAATGG CTAGAGGCCCA TCTTCTCCCAC TCTGAATGG ATTTTCCAAT AGCTCTTCCA TGGTGACTT CTTGAATGA TCTGGTGACTT CTCAAAGA TCTCCGTCCA TCTGGTGACTT CTCAAAGA CCACGTTCCAAA CCACGTTGGG	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACCCTC CATATCCCA AATATCCCA AATATCCCA CCTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TATTGTTG TACAGAGTGC GACTGAATCA TCTGGAAATCA TCGGAAAAAA CAAACTGCGC CAGCCTTGGT GACCTGGTAC CAAGCTGGTAC CAACATTGGAAATT GGATCTCAC CTACTCCCA CTTCGTCCCC CACCTTTCT CATGGTACC CTTCGTCCCC CACCTTTCT CATGGTCCCC CACCTTTCT CATGGTCCCC	TTATTGATCC CTGATGAGTGT GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTTC CTCCCTGC CAATTTTCTG TGCTGTCC CACTAGGCAAA CTTTTAAAAT AGAGGGACAG AATCTGTCA AAAACTTGTT AGGCAGAGC AAGAATGAAG CCTGCCAGCAG CCTGCCAGCAG CCTGCCAGCAG CCTGCCAGCAG CCTGCCAGCAG CCTGCCAGCAG AGAATGAAG CCTCTCTCTA ACCAGCAGCAG ACGAACCC CGGAACTC GAAAATTTCT GACGAGGTGA ACCAGCTGA ACCAGCTGA ACCAGCTGC AGCAGCTC CGAAAATTTCT TCTCCGTGA CCCCGGACCT CCTCAGGACTC CCTCAGGACT CCTCAGGACTC CCTCAGGACT CCTCAGGACTC CCTCAGGACT CCTCAGACT CCTCAGGACT CCTCAGGACT CCTCAGACT CCTCAGGACT CCTCAGGACT CCTCAGC CCTCAGACT CCTCAGGACT CCTCAGGACT CCTCAGGACT CCTCAGGACT CCTCAGACT CCTCAGGACT CCTCAGGACT CCTCAGGACT CCTCAGGACT CCTCAGACT CCTCAGC CCTCAGACT CCTCAGGACT CCTCAGC CCTCAGACT CCTCAGC CCTCACC CCTCAGC CCTCAGC CCTCAGC CCTCAGC CCTCAGC CCTCAGC	TCCAGTCGAG GTCAAAAAAT ACCTCAGTAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTG GTCACTCTG CATAGACTCTGG CTAGGTCTTG CATAGAAAGT TGTCTTATAC TTCATTTTC CCTCTTCCCG CTGAGGAAA CCTCTCGGACA CCTCTGGCAT TCGTTACAA CGATCCCGG CCAACCAGAT TTGATCCTGTACAA TCGTTACAA CCACCAGAT TTGATCCTGC TAGCAT TCGTTACAA CCACCAGAT TCGTTACAA CCACCAGAT TCACTTTCCCG TACTTTCCCG CCACCAGAT TCGTTACAA CCACCAGAT TCACTTCCCG TACTTCCCG TACTTCCCG TACTTCCCC TCCCGGTGT TCCCCGCGTGT	AACATGTATA GACCACAGCG CCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGCCG TTGGTGCTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATTA CTGTCCCAA GTTTGCAATG TCTGTCCCAA GTTTGAATGT TGATGTTGC CAAAGACACT CACTGTGGTC GGAACACTT CACCTTTTT GGTCCGCAT CACTTTTT GGTCCGCATC ATGCTCACT CACGTTTTC CACGTTTTT CACGTCTTTT CAGGCGGTTT CCCTGAGTT CCCTGGTT CCCCGGTTT CCCCTGTTT CCCCTGTTT CCCCTCTTTT CTCCTGTTT CTCCTCTGTT	120 180 240 300 360 420 540 600 660 720 780 840 900 1020 1140 1260 1320 1380 1560 1560 1680 1740 1860 1740 1860 1920
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	TCCAAAGTTC CACAGAAGTTC CACAGAAGTAG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTT TCAGGCTGAC AAATGCCCTC TTTTTTTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT TCTCTCTGTGG AGCCTCCTTC AAAACGCAA TCCTCTCAGC TTGAGGCCAA TTCATTCAC AATAAAAGAA TTTAACTAGG CTGGGGCCCC CTGGGAGCAG ATTATGACTAG TCTCCCTCAA TCTTCTCCCA TCTGAATGG AGAAGACCCA TCTTGAATGG AGAAGGACCTA ATTATCAAT ATTTCCAT TCTGAATGG TCTAGATGT GATTTCCAAT ATGAGTGTT CCTTTCAAAA TCAAGACAGA TCTTTCAAAA TCAAGACAGA CCACGTTGGG ATTTCTAAAA CCACGTTGGG ATTTCTAATT	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA ACTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCT TTTTGTTG TACAGAGTGC GACTGAATCA CAAGCTGATACAAT AGTTGGCAAA CGAGCTTGGT CAAGTGGGC CAGCCTTGGT GACTGGTTAC CAAGTGGGC CAGCCTTGGT GGACATGAG GTACACAATT GGATTCTCAC CTACAGTGG CTTTGATTTT CATGGTCATG CTATGTCCCT GTCTCCTCCC GCCATCTGC CGCCATCTGC CCCTTTCTC	TTATTGATCC CTGATGAGTI TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG ACTAGGCAAA CTTTTAAAAT AGAGGGACAA AATCTGTTA AGAGTGAGA AAACTTGTT AGGCAGAGG CAGCAGTCTCTCTCA CAGCAGAGG CCTCCTCTA ACAGCAGAGC ACCAGCATAT ATCACCATGC AGCAGAGCC AGCAGCTAT ATCACCATGC AGCAGGAGTGA ACGATTTCTCTCTCTCGGAAAATTTCA ACAGGAGTGA ACGATTTTCTTCTCTCGTGAACTTTCTTCCGTGAACTTTCTTCCGTGAACTTTCTTCCGTGAACTTTTCTTCTTCTTCTTTCT	TCCAGTCGAG GTCAAAAAAT ACCTCAGTAG ACTCCTGTTG ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG TCACTCCTGG CCAGCCAGC CTAGGTCTTG CATAGAAAG TTGTTTAGAC CCTCTGGAT TTGTTTAGAC CCTCTGGAT TCCTTTTCC CTCTTCTCCG CTGAGGAAAC CCTCTGGCAT TCGGTTTACAA GGATCCGGA TCTGTTACAA GGATCCGGA TCTGTTACAA GGATCCGGA TCTGTTAGAAAT GCAACAAAAA CCAGTTGAAAT GCAACAAGT TCTGTTAGAAT CCACGGT TCTAGGGT TCAGGGT TCAGGGT TCAGGGT TCCACGGT TCCACGGT TCCACGGT TCCCCCCGGT TCCCCCCGGT TCCCCCCCGGT TCCCCCCCGT TCCCCCCCGGT TCCCCCCCGGT TCCCCCCCGGT TCCCCCCCGGT TCCCCCCCGGT TCCCCCCCGGT TCCTCCCCC	AACATGTATA GACCACAGG CCCCCTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAAG ACGAGTATTA GGAGAGGCCG TTGGTGGTGG GCTCAGACAT GTTTGTTGA TTTGTTGA TTTGTTGA TTTGTTGA GTTTGATATA TCTGTCCCAA GTTTGAACACT CCTGAACACT CCTGAACACT CACTGTTGTT CACCTTTTT GGTCCGCAT CACTGTTTC TAGCTCACTC TAGCTTTTC CAGCTTTTC CAGCTGTTC CAGGGGTTT CCAGGGGTTT CCCTGGGTT CCACCTCTTTT CTCCTGGTT CACCTCTTTT CTCCTGGGTT CCACCTCTTTT CTCCTGGGTT CCCTGTTTCCTCTTTT CTCCTGTTTCCTTTT CTCTTTTTCTTTT	120 180 240 300 360 420 540 600 780 840 900 960 1020 1140 1220 1380 1440 1500 1620 1680 1680 1740 1860 1980 2040
50 55 60 65	TCCAAAGTTC CACAGAAGTTC CACAGAAAGTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC GCTGCTCTTT CACCACCACC CTCTGGCTTT TCAGGCTGAT AAATGCCCTC TTTTTTTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT AGCCTCTTC AAAACGCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAGAG TTAAAGAAAT TTTAACTAGG CTGGGGCCCC AGGGACCTCA ACCCCAGCC CTGGGAGCAG ATTATGACTAC TCTCATCCAC TCTCAATG ACAAGGTGTT AGCAGCC TCTGGAATGG AGAAGGTGTT AGCAGCTCTCAA TTGGAATGG TAGAGGACCTCA TCTGAATGG AGAAGGTGTT ATGAGATGAT ATGAGATGAT TCGTTCAAT TCGTTCAAT TCGTTCAAT TCGTGACTT CCTTTCAAAA TCAAGACAGA CCACGTTGGG ATTTTCAATAT TCAACTTCCT	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG CCTGCCCTC GGGACCAAC GTGTAACATT TGATCATACA TCACAAGCCT TTTTGTTT TACAGAGTGC GACTGAAAAT AGTTGGCAAA CAAACTGCGC CAGCCTTGGCCTC GACTGATCAC CAATGTGGTAA CAAACTGCGC CAGCCTTGGT CCAGCATGAG GTACACAAT CCTACAGTGG GTACACATT CCTACAGTGG CTTACACAC CTTCCCC GTCTGCTCCA CACCTTTCCT CTCTCCCC GATCTTCCC GATCTTCCC	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG TGCCTGCTC ACTAGGCAAA CTTTTAAAAT AGAGGACAG AATCTGTCA CAGATGAC CAGCAGAAG CAGCTCCTCTCTCTCT AGCCAGAAG ACGATGAC AGCAACCA ACGAGTGA ACGATTTCC GAAAATTTCA ACAGGAGGA ACGATTTCCTCTCTCT GAAAATTTCA CAGGAGGCAC CCTCGCAGAAG CCTGCCAGAAG CTTTCCCTCG GAAATTTCA ACAGGAGTGA ACGATTTCT TCTTCCGTGA GCCCGGACCT CGTAGAAAAG	TCCAGTCGAG GTCAAAAAAT ACTCCAGTAG ACTCCTGTTG ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG GTCATCCTGTG CATAGAAAGT TTGTTTATA TTGGTTTTAGAC CTAGGTCTTG CATAGAAAGT TTGATTTTC CTCTCTCCCG CTAGGAAAACT CCAACCAGAT TTGATCCCGGAT TCGTTACAA GGATCCCGA CCAACCAGAT TTGATGCCGG TATCTTTCTC AGCTTGAAAA CCAACAAAAC TCAATGTGAG TCAACAAAAC TCAATGTGAG TCAACGAGT TCCAGGATGCT CCTCAGGAT TCCCCCGGTGT CCCCCGGGT CCTCAGGCTT CCCCCGTGT CCCCATGCTTC	AACATGTATA GACCACAGC CCCCCTITGT GAGTTICATA ATCTTCTTCT GGCAGCCAGT ACCACACAG ACGAGTATTA GGAGAGGCCG TTGGTGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGGTATA TCTGTCCCAA GTTTGATGT CACTGTGGTC CGAACACT CACTGTGGTC CGAACACT CACTGTGGTC CGAACACT CACTGTGGTC CGAACACT CACTGTTTT CACTTTTT GGTCCGCAT ATGCTCACT CAGGGGTT CACGGGTT CACGGGTT CACGCGTT CACCTCGTT CCCTGAGTT CCCTGAGTT CCCTGGGTT CCCTGGGTT CCCCTGTT CCCTCTGTT CCCTCTTTTC CCCTTTTTC CTCTTTTT CTCCTTTTT CTCCTTTTT CTCTTTTT CTCTTTTT CTCTTTTTT	120 180 240 300 360 420 540 600 660 720 780 840 900 1020 1140 1200 1320 1380 1440 1560 1560 1680 1740 1860 1860 1920 1980 2040
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	TCCAAAGTTC CACAGAAGTAC CACTAGAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTTCTTA CAGCAAGGTT CCAGTCACAA GAGGCACTAG ATCCTGAGTA ACTGTTGAGA ATTGCCCCC GAGACTGAGA ATTGACATCA AGGTATCTTTG AGGAATTCTA TACAAGGATG CACATGCTCA AACTCCTGAGA AACTCCTGGA ATTGCCTCTTG AGGAATTCTT TACAAGGATG CACATGCTCTTG ACAACCCAGG ACAACCCTAGA ACTCCTTGGA ACTCCTTGGA ACACCCTGGA ACACCCTTGG CTGACCATGAC CTGACCATGAC CTGACCATGAC CTGACCATGAC CTGACCATGAC CTGACCATGAC CTGACCATGAC CTGACCATCTCT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT AGCCTCCTTC AAAACCGCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAGAAG TTAAACAAGC CTGGGGCCCC AGGCACCTCCAGCC CTGGGAGCAC ATTCTCCCCA TTCTCCCCA TTCTCCCCA TCTCAATCACAC AGCAGCTCTA AGCTCTCAAT AGCTCTTCCAT ATGAGATGAT AGCTCTTCCAAA ACCTCTCAAAA CTTTCAATA CCTAGAAGAGAT TTGAGATGAT AGCTCTTCCAAAA CTTTCAAAA CTTTCAAAA CTTCCAAAA CTTTCAAAA CCACGTTGGG ATTTTCAATA TCAAACAGACAGA CCACGTTGGG ATTTCTAATA TCAAACATCCT GTATCAATAG	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTGTT GGCTGTGGAG CCTCATCCC AATATCCCA ATATTCCCA ATATTCCCA CCTGGCCCTC GGCGACCAAC GTGTAACAT TGATCATAAA TCACAAGCCT TTTTGTGTTG TACAGAGTGC GACTGAAAA CAAACTGCGC CAGCCTTGGT CCAATGTGGC GACCTAGGT CCAATGTGGC GACCTGGTC CAATGTGGCC CAATGTGGCC CAATGTGGCC CAATGTGGCC CAATGTGCCC CAATGTCCC CTACAAGTGG CTTACAAGTGG CTTACAAGTGG CTTAGATCTCC CACCATCTCC CACCATCTCC CACCATCTCC CACCATCTCC CACCATCTCC CACCATCTCC CACCATCTCC CACTGCCCAT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAAGTTG CCTGCCTTCC CAATTTTCTG TGCCTGCC CAATTTTCTG TGCCTGCC ACTAGGCAAA CTTTTAAAAT AGAGGGACAG AAACTTGTCA AAAACTTGTT AGGCAGAGC CAGCAGCAT CCTCCCTCT CTGCCAGAAG CCTGCCTCTCT ACAGAACGCC AGCAGCTAT ATCACCATGC AGCAGCTAT ATCACCATGC AGCAGTGT TTCTCTGTA ACAGGAGTGA ACAGTATTCA ACAGGAGTGA TCTTCCGTC TCTTCCGTC TCTCCTGTAAAATTTCA ACAGGAGTGA ACAGTATTTC TCTTCCGTCA TCTCCGGACCT CGTAAGAAT TTCGCTCTC CGTAAGAAT TTCGCTTACCCC	TCCAGTCGAG GTCAAAAAAT ACTCAGAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG GTCATCCTGGT CATGGTCTTG CATAGAAAGT TTGTTTATC CTTTACTA TTGGTTTTCC CTGAGGAAGC CTCTCGGAT GAGAGAAGC TCCTAGACAT CTGTTACAG CCACCAGAT TCTGTACAT TCTGTACAT TCTGTACAT TCTGTACAT TCTGTACAT TCTGTACAGA CCAACCAGAT TTGATCCGG TATCTTCTC TCTGCAGT TCTGTACAGA CCAACCAGAT TCTGTACAGA CCAACCAGAT TCTGTACAGA CCACGAGT TCTGTAGAAT GCAACAAAAC TCAATGTGA CCACGGTGT CCTCAGGGTT CCCCGGTGT CCTCAGGGTT CCCCAGGTTC CCCCATGCTTC GTCCACGTTC	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTG GGAGCAGTATTA GGAGAGCAGT ACCACAAAG ACGAGTATTA GGAGAGGCGG TTGGTGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGAATGT TGATGTTGC CAAGTCACT CACTGTTGGT CGACACTT CACCTTTTT GGTCCGCAT CACTTTTT CTCGGCATC CAAGGAGAGG TGAATACACC CAAGAGAGG TGAATACACT CACGCATT CACCTTTTT CTCCTATATC CAGGCGGTT CACCTCTTT CTCCTGGGTT CACCTCTTT CTCTGTTGGG TCCTAAACTC CCGAGCCCTGT CCCGAGCCCTGT CCCGAGCCCTGT CCCGAGCCTGT CCCGAGCCCTGT CCCGAGCCCTGT CCCGAGCCCTGT CCCGAGCCCTGT CCCGAGCCCTGT CCCGAGCCCTGT CCCGAGCCCTGT	120 180 240 300 360 420 600 660 720 780 840 900 1020 1140 1260 1320 1380 1440 1500 1620 1680 1740 1880 1740 1890 1980 2040 2160
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	TCCAAAGTTC CACAGAAGTAC GTGTAAAGAG GTGTAAAGAG CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGCT TTTTTCTTA CAGCAAGGTT CCAGTCACCAA GAGGCACTAG TCTAGGCTCC GAGACTGAGA ATTGGCCCC GAGACTGAGA ATTGACATCA ACTGTTGACA TACAAGGAT CACATGCTCA TACAAGGAT TCCTGAGT ACTGTTGACA TACAAGGAT TCCTGGGA ACTCCTGG ACACCCCCAG CCTTTTTGCTTG ACACCCTGG ACACCCTGG TTTTTTGCTTGG ACACCCTTGG TCTTTTTGCTTGG ACACCCTTGG TTTTTTGCTTGG ACACCCTTGG TTTTTTGCTTGG CCCCACCCAAC	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC AGCCAAATCAC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGACCCCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAGAGT TTTAAACTAGG CTGGGGCCCC AGGCCCCAGCCC AGCCCCAGCCC ATTCTTCCCAA TCTTCTCCCA TTCTCTCCCA TCTTCTCCCA TCTTCAAGAGTTT ATGAGTGTT CAAGTTTCCAAT AGCACTTCCA TCTGAATGG ATTTCCAAT AGCACTTCCA TCTGAATGG ATTTCAAAA CCACGTTGGA TCAAACTCCT TCAAACTTCCT TCAACTTCCT TTAACTACA ATCAAGAAGAC ATTCAAAAA ATCAAGAAGAC ATTCAAAAAACACAGAAAGACAGAACACACGTTGGG ATTCTATATT TCAACTTCCT TGTAACTACAACACACACACACACACACACACACACACAC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACCCAC CATGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTTTG TACAGAGTGC GACTGAATCA TCAGAGTGC GACTGAATCA CAACTGCGC CAGCCTTGGT GACCTGGTAC CAACTGGTAC CAACTGCGC CAGCCTTGGT GACCAACTGCGC CAGCCTTGGT GACTACAATT CATGGTAC CTACAAGTGC CTACACATCT CATGCTCCC GTCTCCC GCCATCTCC GACCATCTCC GACCATCTGC CATCTGCCAC TTTTTTTTTT	TTATTGATCC CTGATGAGTGT GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CATATTTCTG TGCCTGCC CAATTTTCTG TGCCTGCC CAATTTCTG TGCCTGCC CATAGGCAAA CTTTTAAAAT AGAGGGACAG AATACTGTTA AGAGCAGAGC	TCCAGTCGAG GTCAAAAAAT ACCTCAGTAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCATCTGGT TCACTCTGG GTCATCTGG CTAGGTCTTG CATAGAAAGT TGTCTTATAC TTCATTTTC CCTCTTCCG CTGAGGAAA CCTCTCGACAT CCTCTGAGCAAT CCTCTTAGACA TCGTTACAA CCACCAGAT TTGATCCGG CTAGCAGAT TCATTTTCA GCACAGAT TCAGTTACAA CCACCAGGT TTCACTCCGG TACTTTCCG CCACCAGCT TCCCGGGTG CCCCAGCTT CCCCAGCTT CCCAGCGTT CCCACGTTC CCCATGCTT CCCACGTTC CCCATGCTT CCCACGTTC CCCATGCTT CCCACGTTC CCACTTCACGGGGG	AACATGTATA GACCACAGCG CCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGCCG TTGGTGCTGG GCTCAGACAT GTTTGTTGA CCCCAAAGA GTTTGCTATTA CTGTCCCAA GTTTGCAATG TCTGTCCCAA GTTTGAATGT TGATGTTGC CAAAGACACT CACTGTGGTC GGAACACTT CACCTTTTT GGTCCGCATC ATGCTCACT CACTTTTT GGTCCGCATC ATGCTCTCTAACTC CAAGGGGGTTT CTCCTGAGTT CTCCTGGTT CCCTGATATC TCCCTATATC TCCCTATATC TCCCTATATC TCCCTAGTGAGAG TCCTAAACTC TCCCTATATC TCCCTATATC TCCCTATATC TCCCTAGTGAGAG TCCCAAGGGGGTT CCCCAAGGGGGTT CCCCAAGGAGAG TCCCTATATC TCCCTATATC TCCCTATATC TCCCTAGAGGAG TCCCAAGGGGGTT AAGTGATGGA	120 180 240 300 360 420 540 600 660 720 780 840 900 1020 1140 1200 1320 1380 1440 1560 1560 1680 1740 1860 1860 1920 1980 2040
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	TCCAAAGTTC CACAGAAGTAC GTGTAAAGAG GTGTAAAGAG CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGCT TTTTTCTTA CAGCAAGGTT CCAGTCACCAA GAGGCACTAG TCTAGGCTCC GAGACTGAGA ATTGGCCCC GAGACTGAGA ATTGACATCA ACTGTTGACA TACAAGGAT CACATGCTCA TACAAGGAT TCCTGAGT ACTGTTGACA TACAAGGAT TCCTGGGA ACTCCTGG ACACCCCCAG CCTTTTTGCTTG ACACCCTGG ACACCCTGG TTTTTTGCTTGG ACACCCTTGG TCTTTTTGCTTGG ACACCCTTGG TTTTTTGCTTGG ACACCCTTGG TTTTTTGCTTGG CCCCACCCAAC	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC AGCCAAATCAC TGGCCTTTAA TGTGGTGCAT CTGTCTGTGG AGACCCCAA TCCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAGAGT TTTAAACTAGG CTGGGGCCCC AGGCCCCAGCCC AGCCCCAGCCC ATTCTTCCCAA TCTTCTCCCA TTCTCTCCCA TCTTCTCCCA TCTTCAAGAGTTT ATGAGTGTT CAAGTTTCCAAT AGCACTTCCA TCTGAATGG ATTTCCAAT AGCACTTCCA TCTGAATGG ATTTCAAAA CCACGTTGGA TCAAACTCCT TCAAACTTCCT TCAACTTCCT TTAACTACA ATCAAGAAGAC ATTCAAAAA ATCAAGAAGAC ATTCAAAAAACACAGAAAGACAGAACACACGTTGGG ATTCTATATT TCAACTTCCT TGTAACTACAACACACACACACACACACACACACACACAC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACCCAC CATGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTTTG TACAGAGTGC GACTGAATCA TCAGAGTGC GACTGAATCA CAACTGCGC CAGCCTTGGT GACCTGGTAC CAACTGGTAC CAACTGCGC CAGCCTTGGT GACCAACTGCGC CAGCCTTGGT GACTACAATT CATGGTAC CTACAAGTGC CTACACATC CTACACTCCC CTCCCCC CCCCTTCCC GCCATCTCC GACCCTTTCT CCCCACTCTCC CACCCTTTCC CGCCATCTGC CATCTGCCCAC TTTTTGTGTGC	TTATTGATCC CTGATGAGTGT GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CATATTTCTG TGCCTGCC CAATTTTCTG TGCCTGCC CAATTTCTG TGCCTGCC CATAGGCAAA CTTTTAAAAT AGAGGGACAG AATACTGTTA AGAGCAGAGC	TCCAGTCGAG GTCAAAAAAT ACCTCAGTAG CACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCATCTGGT TCACTCTGG GTCATCTGG CTAGGTCTTG CATAGAAAGT TGTCTTATAC TTCATTTTC CCTCTTCCG CTGAGGAAA CCTCTCGACAT CCTCTGAGCAAT CCTCTTAGACA TCGTTACAA CCACCAGAT TTGATCCGG CTAGCAGAT TCATTTTCAATTTCC TCAGCAGAT TCGTTACAA CCACCAGAT TCGTTACAA CCACCAGAT TCAGTTACAA CCACCAGAT TCAGTACAA CCACCAGAT TCAGTACAC TCACGATGGT CCCCAGCGTT CCCCAGCGTT CCCACGTTC CCCATGCTTC CCACTGAGGGT CCCCATGCTTC CCACTGAGGGT CCCCATGCTTC CCCACTGTTC CCACTGAGGGT CCCCATGCTTC CCACTGAGGGT CCCCATGCTTC CCACTGAGGGT CCCCATGCTTC CCACTTCACCGGTT CTCCCACGTTC CCACTTCACCAGGGGT CCCCATGCTTC CCACTTCACCAGGGGT CCCCATGCTTC CCACTTCACCAGGGGT CCCCATGCTTC CCACTTCACCAGGGGGT CCCCATGCTTC CCACTTCACCAGTTC CCACTTCACCAGTTC CCACTTCACCAGTTC CCACTTCACCAGGGGT CCCACTTCACCAGGGGT CCCACTTCACCAGGGGT CCCACTTCACCAGGGGT CCCACTTCACCAGGGGT CCACTTCACCAGGGGT CCCACTTCACCAGGGGT CCCACTTCACCAGGGGT CCCACTTCACCAGGGT CCACTTCACCAGGGGT CCACTTCACCAGGGT CCACTTCACCAGGGT CCACTTCACCAGGGT CCACTTCACCAGGGT CCACTTCACCAGGGT CCACTTCACCAGGGT CCACTTCACCAGGGT CCACTTCACCAGGGGT CCACTTCACCAGGGGT CCACTTCACCAGGGGT CCACTTCACCAGGGGT CCACTTCACCAGGGGT CCACTTCACCAGGGGT CCACTTCACCAGGGT CCACTTCACCAGGGT CCACTTCACCAGGGT CCACTTCACCAGGGT CCACTTCACCAGGGT CCACTTCACCAGGGT CCACTTCACCAGGT CCACTTCACAGGT CCACTTCACCAGGT CCACTTCACAGGT CCACTTCACAGG CCACTTCACAGG CCACTTCACAGG CCACTTCACAGG CCACTTCACAGG CCACTTCACAGGT CCACTTCACAGG CCAC	AACATGTATA GACCACAGCG CCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGAGCCG TTGGTGCTGG GCTCAGACAT GTTTGTTGA CCCCAAAGA GTTTGCTATTA CTGTCCCAA GTTTGCAATG TCTGTCCCAA GTTTGAATGT TGATGTTGC CAAAGACACT CACTGTGGTC GGAACACTT CACCTTTTT GGTCCGCATC ATGCTCACT CACTTTTT GGTCCGCATC ATGCTCTCTAACTC CAAGGGGGTTT CTCCTGAGTT CTCCTGGTT CCCTGATATC TCCCTATATC TCCCTATATC TCCCTATATC TCCCTAGTGAGAG TCCTAAACTC TCCCTATATC TCCCTATATC TCCCTATATC TCCCTAGTGAGAG TCCCAAGGGGGTT CCCCAAGGGGGTT CCCCAAGGAGAG TCCCTATATC TCCCTATATC TCCCTATATC TCCCTAGAGGAG TCCCAAGGGGGTT AAGTGATGGA	120 180 240 300 360 420 600 660 720 780 840 900 1020 1140 1260 1320 1380 1440 1500 1620 1680 1740 1880 1740 1890 1980 2040 2160
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	TCCAAAGTTC CACAGAAGTTC CACAGAAAGTTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAC AAATGCCCTC TTTTTTTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAA TCTCTCTGTGG AGCCTCCTTC AAAACGCAA TCCTCTCAGC TTGAGGCCAA TTCATTTCAC AATAAAAGAG TTAAACAAGA TTTAACTAGG CTGGGGCCCC AGGGACCTCC CTGGGAGCAG ATTATGACCA TCTCCGTCAA TCTTCCCCA TCTGAATGG AGAAGGTGTT AGCAAGGTGTT CATTCCAAT ATGAGATGAT TTGAGACTGA TTGGAGACTCCA TCTGAATGG AGAAGGACTCAA TCTGCTCAAT AGCACGTTGGG CCACGTTGGG CCACGTTGGG ATTTCTAAAA TCAAGACAGA ATCAGGAAGC CCATTTCTATT TCAACTTCCT GTATCAATAG ATCAGGAAGC CCGTCTTGCTC CCTTTCCAATA TCAACTTCCT GTATCAATAG ATCAGGAAGC CCGTCTTGCTC CCTTTCCATCCT CCTTTCCAATAG CCACGTTGGG CCTCTTCCTCC GTATCAATAG ATCAGGAAGC CCGTCTTGCTC CCTTTCCTCCTC CCTTTCCTCCTC CCTTTCCTCC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG CTGCCTC CCTCATCCC AATATTCCCA ACTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTGTT TACAGAGTCC GACTGAATCA CAACTGGACAA CGAGCTTGGT CAACTGGTCA CAACTGGGC CAGCCTTGGT GACCTGGTT CCAACTGGT CCAACTGGT CCAACTTGCT CCACCTTCTC CTATGCCCT GTCTGCTCA CCCTTTTCT CCCCATCTGC GACCTTGCT CCCCTTCTCT CCCCATCTGC GATCTACAAC CCCTTCCTCC CACCTTTTCT CTCCCCCACCTTTCT CCCCATCTGC GATCTACAAC CCCTTCCCC CACCTTTCT CCCCATCTGC GATCTACAAC CCCTTCCCC CACCTTTCT CCCCACCACAC AGCCCAGCAG AGCCAGCAG AGCCCAGCAG AGCCAGCAG AGCCCAGCAG AGCCAGCAG AGCCCAGCAG AGCCAGCAG AGCCAGCAGAG AGCCAGCAG AGCCAGCAG AGCCAGCAG AGCCAGCAG AGCCAGCAG AGCCAGCAG AG	TTATTGATCC CTGATGAGTI TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG TGCCTGCTCC ACTAGGCAAA CTTTAAAAT AGAGGGACAA AAACTTGTT AGGCAGAGA CAGCTCCTCT CTGCCAGAAG CCTGCTCTCTA ACAGAGTGAC AGCAGACTCT ACAGGAGTAC ACAGGAGTAC TCTTCCCTG GAAAATTTCT CAGAGAGTCCT CTCCCGGACT TCTTCCGTGA TCTTCCCTC GAAAATTTCT TCTTCCGTGA CCCCGGACT TTCGTCTCTC CGTAAGAATT TCGTTCTCT CGGACAAAAG GCCCGTACCC CCCGTACCC CCCCTAGCC	TCCAGTCGAG GTCAAAAAAT ACCTCAGTAG ACTCCTGTTG ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG GTCATCCTGG CTAGGTCTTG CATAGAAAG TTGTTTAGAC CTAGGTCTTG CATAGAAAG TTGTTTTTC CTCTTTATAA TTGGTTTGCT TTCATTTTC CTCTTCTCGCAT GAGAGAAC CCTCTGGCAT TCTGTTACAA GGATCCGGA TCTGTAGAAT TCAGTTTACAA GGATCCGGA TCTTGAAAAT TCAGTTTACA GCACAGAGT TCTGTTACCG TCTTAGAAAT CCACGGTT CCACGGTT CCACGGTT CCACGGTT CCCCCGGTT CCTCGCGC CCCATGCTTC GTCCACGTTC CCCCCGTTC CCCATGCTC CCCATGCTC CCCATGCTC CCCATGCCC CCATGCTCC CCACTAGCCC CCATGCTACCC CCATGCTAGCCC CCATGCTAGCCC CCATGCTAGCCC	AACATGTATA GACCACAGGG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAAG ACGACTATTA GGAGAGGCCG TTGGTGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGGA TTTGGTATA TCTGTCCCAA GTTTGAATGT CAAGTGTTGTC GGAATCACT CACTGTGGTC CGGAATCACC TAGCTTTTC CACTTTTT GGTCCGCATC ATGCTCACT TAGCTTTTC CAGCGGTTT CACCTTTTC CAGGGGTT CCCGGGTT CCCGGGTT CCCTATAT CTCTCTATAT CTCTTTGGT CTCCTATAT CTCTTTTGGT CAGCGGTTT CACCTCTTTT CACCTCTTTT CACCTCTTTT CACCTCTTTT CACCTCTTTT CACCTCTTTT CACCTCTTT CACCTCTTTT CACCTCTTT CACCTCTT CACCTCT CACCTCT CACCTCT CACCTCT CACCTCT CACCTCT CACCTC CACCTCT CACCTCT CACCTCT CACCTCT CACCTC CACCT CACCTC CACCT CACCTC CACCT CACCTC CACCT C	120 180 240 300 360 420 600 780 840 900 960 1020 1140 1200 1140 1250 1380 1440 1500 1680 1740 1860 1920 1980 2160 2220 2280
50 55 60 65 70	TCCAAAGTTC CACAGAAGTTC CACAGAAAGTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGAT AAATGCCCTC TTTTTTTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAT AGCCTCTTC AAAACGCAA TCCTCTCAGC TTGAGGCCAA TCATTTCAC AATAAAAGAG TTAAAGAAAT TTTAACTAGG CTGGGGCCCC AGGGACCTCA ACCCCAGCC CTGGGAGCAG ATTATGACTAC TCTCATCAAT TCTTCACAAT TCTTCACAAT ACCTCTCAA TCGTGCAA TCGTGAATGG AGAAGGTGTT AGAGTAGT AGCAGTTCCA TTGGTGACTT CCTTTCAAAA TCAAGAAGAC TCTCTCAAT TCAACTTCCT CCTTTCAAAA TCAAGACAGA CCACGTTGGG ATTATAT TCAACTTCCT GTATCAATAG ATCAGAAAGC CCGTCTTGCTC GTATCAATAG ACCAGAGGAGC CCGTCTTGCTC GTATCAAAAC CCGTCTTGCTC CGTCTAGCTC CGTCTAGCTC CGTCTAGCTC CGTCTCAAAC TCAAGAAGC CCGTCTTGCTC CGTCTCAAAC TCAACACACC CCGTCTTGCTC CGTCTCAAACC CCGTCTTGCTC CGTCTCAAACC CCGTCTTGCTC CGTCTCAAACC CCGTCTTGCTC CGTCTCAACCC CGTCTTGCTC CGTCTCAACCCT CGTCTTCCTC CGTCTCCAACCCT CGTCTTCCTC CGTCTCCAACCCT CTCTCCAACCCT CTCTCCAACCCT CTCTCTCCTC CGTCTCAACCCT CTCTCTCAACCCT CTCTCTCCTC CGTCTCAACCCT CTCTCTCTCTC CGTCTCAACCCT CTCTCTCAACCCT CTCTCTCTCC CGTCTCAACCCT CTCTCTCAACCCT CTCTCCAACCCT CTCTCTCCTC CGTCTCCAACCCT CTCTCTCCAACCCT CTCTCCAACCCT CTCTCTCCAACCCT CTCTCCAACCCT CTCTCTCCTC CGTCTCAACCCT CTCTCTCCAACCCT CTCTCTCCAACCCT CTCTCCAACCCT CTCTCCAACCCT CTCTCCAACCCT CTCCAACCCT CTCCAACCC CTCCAACCCT CTCCAACCCT CTCCAACCCT CTCCAACCCT CTCCAACCCT CTCCAACCT CTCCAACCCT CTCCAACCCT CTCCAACCCT CTCCAACCCT CTCCAACCCT CTCCAACCC CTCCTCACCT CTCCAACCCT CTCCAACCC CTCCTCAACCC CTCCTCAACCC CTCCAACCC CTCCTCAACCC CTCCTCAACCC CTCCTCAACCC CTCCTCAACCC CTCCTCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACC CTCCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC CTCCTCCAACCC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTGTT GGCTGTGGAG CCTGCCTC GGGACCAAC GTGTAACATT TGATCATACA TCACAAGCCT TTTTGTTT TACAGAGTGC GACTGAACA TCTGGAAAAT AGTTGGCAAA CAAACTGCGC CAGCCTTGGCCTC GACCTAGGT CCAGCCTTGGTACAAC CAAACTGCGC CAGCCTTGGT CCAGCATGAG GTACACAATC CCAGCATGAG GTACACAAT CCACCTTTCT CATGGTCAC CTACAGTGG GTTTGATTT CATGGTCAC CTACATTCCC GACCTTGCC GACCTTGCC GACCTTGCC CACCTTTCCT CGCCATTTCCT CGCCATTTCCA CCCGTGCCCAT TTTTGTTGTGC AGCCCAGCAG GGCCTGCTGT AGGCCAGCAG GGCCTGCTGT	TTATTGATCC CTGATGAGTT TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG TGCCTGCTC ACTAGGCAAA CTTTTAAAAT AGAGGAACA AATCTGTCA AAACTTGTT AGGCAGAGC CAGCAGAAG CCTCCTCTCTCTCT AGCCAGAAG ATCACCATGC AGCAACGCA ACGATGCA ACGATTTCC GAAAATTTCA ACAGGAGGAA ACGATTTCCTCTCTTCT TCTTCCGTGA ACGATGTCA ACAGAAGGACA CCCCTAGCC CAGATGGCC CAGATGACC CAGATTGTCA CAGCACAAAAG GCCCGTACCC CAGATTGTCA CCCCCTAGCC GAGTGGTGCA CCCCTAGCC GAGTGGTGCA	TCCAGTCGAG GTCAAAAAAT ACTCCAGTGAG ACACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG TCACTCTGG TCACTCTGG TCACTCTGG CTAGGTCTTG CATAGAAAGT TTGTTTACAA TCGGTTTACCAG GTCACCAGAACACAGAT TTGATCTCTCAG CCAACCAGAT TTGATCCGGA CCAACCAGAT TTGATCCCGG TACTCTTCCC ACCTTGCAC TCCTTGCAC TCCACGTGCT CCCACGTGCT CCCCACGTTC CCCACTGCTC CCCACTGCTC CCCACTGCTC CCCACTGCTC CCCACTGCTC CCACTGCTC CACCTAGCCC AGCGTTTTAA	AACATGTATA GACCACAGG CCCCCTTGT GAGTTTCATA ATCTTCTTGT GGCAGCCAGT ACCACACAGG ACGAGTATTA GGAGAGGCCG TTGGTGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATA TCTGTCCCAA GTTTGATGT CCCAAAGAG GTTGGATCACCT CACTGTGGTC CGAACACT CACTGTGGTC CGAACACT CACTGTTTT GGTCCCCAT CACTGTTTT CACCTTTTT CACCTTTTT CACCTTTTT CACCTTTTT CACCTTTTC CAAGAGAG TGAAATACAC CCAGCACCTT CACGGGTT CCCCATGTGTC CCGAGCCTT CCCCTATGT CCCCTATACT CCCGAGCCTGT AAGTGATGGA TCCTAAACTC CCGAGCCTGT AAGTGATGGA TCCTAAACTC CCGAGCCTGT AAGTGATGGA TCCTAAACTC CCGAGCCTGT AAGTGATCGC GAAGTACTC CCGAGCCCCG GAAGTACTC CCAAGTCCCC GAAGTACTC CCAAGTACTC CCGAGCCCCC GAAGTACTC	120 180 240 300 360 420 6600 6600 720 780 840 900 1020 1180 1260 1380 1440 15500 1680 1740 1860 1920 1980 2040 2160 2220 2230 2340
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	TCCAAAGTTC CACAGAAGTTC CACAGAAAGTAC CACTTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTTT CACAACCAAC CTCTGGCTTT TCAGGCTGCT TTTTTCTTA CAGAAGCAAG TCTAGGCTCC TATTTCCTAA GAGGCACTAG TCTAGGGTCG AATCCTGAGTA ACTCTGAGTA ACTCTTGAGAA ATTGACATCA GAGAATTCTA TACAAGGATG CACATGCTCA ACATGCTCC GACACTTGG ACACCCAGG ACACCCTGG TTTTTTGACATCA ACTCTTTTGACA CACTCTTTGACAAC CACTCTTTTTTTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAT AGCCAAATCAC TGTGGTGCAT TGTGGTGCAT TGTGTGTGAA TCCTCTCAGA TCCTCTCAGA TCCATTCAC AATAAAGAG TTAAAGAAG TTAAAGAAG TTAAACAGG AGGACCTCA AGCCCCAGCC CTGGAGCAC ATTCTCCCA TTCTCCCCA TTCTCCCCA TTCTCCCCA TTCTGAATGG AGAGGACCCA GCAAGGTGTT GATTTCCAT ATCAGATGAT ACCCTTCCA TCCGTGAATGG AGAGGACCTA TCTGTGATTG ATTCAATAG CCACGTTGGG ATTTCTAAT CCATTCAATAG CCACGTTCGG ATTCTAATAT CCATCAATAG ATCAGGAAGC CGTTCTCCT CTTCTCAATTGG CTTCTCAATAG CCACGTTCGG CTTCTCAATAG CCACGTTCGCT CTTCTCAATAG ATCAGGAAGC CGTTCTCCT CCTTTCCAATAG ATCAGGAAGC CGTTCTGCTC CCTCCAAGCT CCCCAAGCT CCCCAAGCT CCCCAAGTTGTGA	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTGTT GGCTGTGGAG GTGGAGGAC CCTTCATCCC AATATTCCCA ATATTCCCA CTGGCCCTI GGGGACCAAC GTGTAACAT TGATCATACA TCACAAGCCT TTTTGTGTTG TACAGAGTGC GACTGAAAA CAAACTGCGC CAGCCTTGGT CAACTGGAAAA CAAACTGCGC CAGCCTTGGT GACCTGGTAC CAATGTGGTG GACCTGGTAC CTACAAGTGG GTTCACAATTC CGATCATCC CTACAAGTGG GTTTGATTTT CATGGTCCCA CTTCTACAAGTGG GTTTGATTTT CATGGTCCCA CACCTTTTCT CGCCATCTGC CACCTTTCTC CGCCATCTGC CACCTGCCAT TTTTGTGTGC AGCCCAGCAG AGCCCAGCAG AGCCCAGCAG AGCCCAGCAG AGCCCAGCAG AGCCCAGCAG AGCCCAGCAG AGCCCAGCAG AGCCCAGCTGC GGCCTGCTGC GGCCAGTACC GGGCCTGCTGG GGCCTGCTGG GGCCAGTACC	TTATTGATCC CTGATGAGTTG GGACTTCTAA TAATTCAAGTTG CCTGCCTGC CAATTTTCTG TGCCTGCC CAATTTTCTG TGCCTGCC CATTTCTCC CATAGGCAAA CTTTTAAAAT AGAGGGACAG AATCTGTCA AAAACTTGTT AGGCAGAG CAGCTCCTCT CTGCCAGAAG CCTGCCTCTCT AGCGACAGC AGCAGCTAT ATCACCATGC AGCAGGTAT ATCACCATGC AGCAGGTGTA ACAGATTTCC TCTCCTGTG ACAGATTTCC TCTCCTGTG ACAGATTTCC TCTCCTGTGA ACAGATTTCC TCTCCTGTGA ACAGATTTCC TCTCCTGTGA ACAGATTTCC TCTCCTGTGA ACCAGTTTCCCTC CGAAAATTTCC CCCCTACCC CAGATTGTCA CCCCTAGCC CGGATGGCA TGGCAGCGG	TCCAGTCGAG GTCAAAAAAT ACCTCAGTGA ACTCCTGTTA ACTCCTGTTGA CTCCTGTTGAGC CCCAGTGCA GTCATCTGG TCACTCTGG CTGGTCTTG CATAGAAGT TTGTTTATT CCTCTTTCCCG TGAGTTTTCCCTGTTCCCGAT GAGAAAGC TCCTTGGCAT CCTCTGGCAA CCACCAGAT CTGTTACAG CCACCAGAT TTGATCCCGG TAGTTCTCCGAT TCGTTACAG CCACCAGAT TCTGTTACAG CCACCAGAT TTGATCCCGG TATCTTCTC ACCTTGACAT TCGTTACAGA CCAACCAGAT TTGATGCCGG TATCTTTCTC ACCTTGAAAT GCAACAAAAC CCACCAGTGT CCCCACGGTT CCCCCGCGTT CCCCCAGTTCCCCACGTTCCCCACGTTCCCCACGTTCCCCACGTTCCCACGTTCCCACGTTCTAAGGGG CCACTGAGGG CCACTGCTCC CCACTGAGGG CCACTGAGGG CCACTGCTCC CCACTGAGGG CCACTGCTCC CCACTGCTCC CCACTGAGGG CCACTGCTCC CCACTGCTCC CCACCTTTTAA	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGGGCCGG TTGGTGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATA TTTGCTATTA TCTGTCCCAA GTTTGAATGT TCAGTCTCCCAA GTTTGAATGT TGATGTTTCC CAAGACACT CACTGTGGTC GGAATACACC CGACACTTT CACCTTTTT GGTCCGCATC ATGCTCATATC CAGGGGGTTT CACCTGTTTC CACTGTGTTC CACTGTGTT CTCCTATATC TCCTGTTGTGT TCCTGTTTC CCGAGCCTGT AGGTGATC CCGAGCCTGT CCGAGCCTGT CCCTATATC CCGAGCCTGT AGGTGTCCC CAAGGGGTCCC CGAGGCCTCT CCGAGCCTGT AGGTGTCCC CAAGGGGTCCC CGAGGCCTCC CAAGTGACACC CCGAGCCTTT CCCGAGCCTGT AGGTGACTCC CCAAGGGTCCC CGAGGCTCC CAAGTGACTCC CATCCATGTC CCTCCTGTTC CCGAGCCTCT CCGAGCCTCT CCGAGCCTCT CCGAGCCTCC CAAGTACTTC CCGAGCCTCC CAAGTACTTC CCAATGTCC CCAAGTCTCC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CACTCCATGTC CACCTCTTC CATCCATGTC CATCCATGTC CATCCATGTC CACCTCTTTT CACCTCTCTC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CACCTCTTC CATCCATGTC CATCCATGT CATCATT CATCCATGT CATCCATGT CATCCATGT CATCCATGT CATCCATGT CATCCATGT	120 180 240 300 360 420 600 660 780 840 900 1020 11200 1260 1320 1380 1500 1620 1680 1740 1860 1740 1860 1920 2160 2210 2220 2230 2240
50 55 60 65 70	TCCAAAGTTC CACAGAAGTTC CACAGAAAGTTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGG GCTGCTCTTT CACACACAC CCTCTGGCTGT TCAGGCTGAC AAATGCCCTC TTTTTTTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCTTTAA TGTGGTGCAT TCTCTCTGTGG AGCCTCCTTC AAAACGCAA TCATTCACC ATTAAATAAAAGAC TTAAACTAGG CTGGGGCCCC CTGGGAGCAC TCTTCCCAA TCTTCTCCCA TCTTCTCCCA TCTTCAACAC TCTTCAACAT TCTAAGATGAT TTTAACTAGC TTCTGAATGCA TCTTCTCCCA TCTTCAAAA TCTAGGTGTT CATTTCAATA AGCTCTTCCAT TCTAAGACAGA TCTTCTCAAAA TCAAGACTGT CCATTGGACTTC CCTTTCAAAA TCAAGACTGAC TCTTCAACTTCCT GTATCAATAG ATCAGGAAGC CCTCTTCCTC GTATCAATAG ATCAAGTCTC GTATCAATAG ATCAGGAAGC CCTCTTGCTC CCTTCCAAGCT CCGATTGTGA ATCAGGAAGC CCTCTCAAGCT CCGATTGTGA ATAACTACTC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTGTT GCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA ACTGGCCTC GGCGACCAAC GTGTAACATT TGATCATAGAGTGC GACTGAACT TTTGTGTTG TACAGAGTGC GACTGAACT AGTTGGCAAA AGTTGGCAAA AGTTGGCAAA CGAGCATGAG GACTGGTAC CAAGCATGGTAC CAAGTGGTG GTACACAATT GGATTCTCAC CTACAGAGTG CTACAGTGG GTTTGATTT CATGGTCAT CATGTCCCT GTCTGCTCCC GTCTGCTCCC GTTTGCTCCC GATTTTCGTTCCC GACTGCCCAT TTTTTGTGTGC AGCCCAGCAG GGCCTGCCCAT	TTATTGATCC CTGATGAGTI TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG ACTAGGCAAA CTTTTAAAAT AGAGGGACAG AAACTTGTTA AGAGGACAG AAACTTGTT AGGGCAGAGG CCTGCCTCT CTGCCAGAAG CCTTCTCTCT GACGAACGC AGCAGCATAT ATCACCATG ACCAGCATT TCTTCCGTGA ACCAGCAGTAT TCTTCCGTGA TCTTCCGTGA TCTTCCGTGA CCGAACAC CCGAACTTCT CCGAACAC CCGAACAC CCGAACAC CCCGAACC CCCGAACC CCCGAACC CCCGAACT TCCCTCCCAGCC CAGATTGTCA CCAGATTGTCA CCCCTAGCC CAGATTGTCA CCCCTAGCC GAGTGGTGA TGCCCCTAGCC GAGTGGTGA TGCCCCTAGCC GAGTGGTGA TGCCCCGAACAT TCCCCCTAGCC GAGTGGTGCA TGCCCAGTAGCT TTCCCAGTGA TTCCCCAGTGGC TTCCCAGTGGA	TCCAGTCGAG GTCAAAAAAT ACTCCTGTTA ACTCCTGTTG CTGTTTAGTA ACTCCTGTTG CTGTTTAGTA ACTCCTGTTG CTGTTTAGAC CCCAGTGCA GTCCTCTGG GCGGCCAGC CTAGGTCTTA TTGTTTATAA TTGTTTTTCC CCTCTTGCAT TCCATTTTCC CCTCTTGCAT TCCATTTTCC CTCTAGACAT TCTGTTACAA GGATCCCGGA TCTGTTACAA GGATCCCGGA TCTTTCTC AGCTTGAAAT CCAGCTTTCTC AGCTTGAAAT GCAACAAAA CCAGTTCCCG CCCAGGTGT CCCAGGTGT CCCATGCTTC GTGCACGTT CCCATGCTTC CTCCAGGGG CCCATGCTTC CTCCAGGGG CCCATGCTTC CCCATGCTTC CCCCTTGGGGG CCCCTTGGCC CCCATGCTTC CCCCTTGGGGC CCCCTTGGGGC CCCCTTGTTCTC GTGCACGTTC CCCCTTGGGGC CCCCTTGTCTCT CTTCTTCTT	AACATGTATA GACCACAGG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAGA ACGAGTATTA GGAGAGGCCG TTGGTGGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGATGTTGT CAAGTCACAT CCTGAACACT CAGACACT CAGACACT CAGACACT CACAGACACT CAGCACCTTT CACCTTTTT CACCTCTTT CACCTCTTT CACCTCTTT CACCTCTTT CACCTCTTT CACCTCTTT CAGGCGGTT CACCTATATC CCGAGCCTGT AAGTGATGGA TCAGGGGTCC CAAGAGTACTC CATCAGTGTC CATCATGTC CATCATGTC CATCATGTC CATCATGTC CATCATGTC CATCATGTC CATCATGTC CATCATGTC CTTCAATGTC	120 180 240 360 420 660 720 780 840 900 1020 1140 1260 1320 1380 1560 1680 1740 1880 1980 2040 2160 2220 2280 2340 2460
50 55 60 65 70	TCCAAAGTTC CACAGAAGTTC CACAGAAAGTTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGG GCTGCTCTTT CACACACAC CCTCTGGCTGT TCAGGCTGAC AAATGCCCTC TTTTTTTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCTTTAA TGTGGTGCAT TCTCTCTGTGG AGCCTCCTTC AAAACGCAA TCATTCACC ATTAAATAAAAGAC TTAAACTAGG CTGGGGCCCC CTGGGAGCAC TCTTCCCAA TCTTCTCCCA TCTTCTCCCA TCTTCAACAC TCTTCAACAT TCTAAGATGAT TTTAACTAGC TTCTGAATGCA TCTTCTCCCA TCTTCAAAA TCTAGGTGTT CATTTCAATA AGCTCTTCCAT TCTAAGACAGA TCTTCTCAAAA TCAAGACTGT CCATTGGACTTC CCTTTCAAAA TCAAGACTGAC TCTTCAACTTCCT GTATCAATAG ATCAGGAAGC CCTCTTCCTC GTATCAATAG ATCAAGTCTC GTATCAATAG ATCAGGAAGC CCTCTTGCTC CCTTCCAAGCT CCGATTGTGA ATCAGGAAGC CCTCTCAAGCT CCGATTGTGA ATAACTACTC	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTGTT GCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA ACTGGCCTC GGCGACCAAC GTGTAACATT TGATCATAGAGTGC GACTGAACT TTTGTGTTG TACAGAGTGC GACTGAACT AGTTGGCAAA AGTTGGCAAA AGTTGGCAAA CGAGCATGAG GACTGGTAC CAAGCATGGTAC CAAGTGGTG GTACACAATT GGATTCTCAC CTACAGAGTG CTACAGTGG GTTTGATTT CATGGTCAT CATGTCCCT GTCTGCTCCC GTCTGCTCCC GTTTGCTCCC GATTTTCGTTCCC GACTGCCCAT TTTTTGTGTGC AGCCCAGCAG GGCCTGCCCAT	TTATTGATCC CTGATGAGTI TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG ACTAGGCAAA CTTTTAAAAT AGAGGGACAG AAACTTGTTA AGAGGACAG AAACTTGTT AGGGCAGAGG CCTGCCTCT CTGCCAGAAG CCTTCTCTCT GACGAACGC AGCAGCATAT ATCACCATG ACCAGCATT TCTTCCGTGA ACCAGCAGTAT TCTTCCGTGA TCTTCCGTGA TCTTCCGTGA CCGAACAC CCGAACTTCT CCGAACAC CCGAACAC CCGAACAC CCCGAACC CCCGAACC CCCGAACC CCCGAACT TCCCTCCCAGCC CAGATTGTCA CCAGATTGTCA CCCCTAGCC CAGATTGTCA CCCCTAGCC GAGTGGTGA TGCCCCTAGCC GAGTGGTGA TGCCCCTAGCC GAGTGGTGA TGCCCCGAACAT TCCCCCTAGCC GAGTGGTGCA TGCCCAGTAGCT TTCCCAGTGA TTCCCCAGTGGC TTCCCAGTGGA	TCCAGTCGAG GTCAAAAAAT ACTCCTGTTA ACTCCTGTTG CTGTTTAGTA ACTCCTGTTG CTGTTTAGTA ACTCCTGTTG CTGTTTAGAC CCCAGTGCA GTCCTCTGG GCGGCCAGC CTAGGTCTTA TTGTTTATAA TTGTTTTTCC CCTCTTGCAT TCCATTTTCC CCTCTTGCAT TCCATTTTCC CTCTAGACAT TCTGTTACAA GGATCCCGGA TCTGTTACAA GGATCCCGGA TCTTTCTC AGCTTGAAAT CCAGCTTTCTC AGCTTGAAAT GCAACAAAA CCAGTTCCCG CCCAGGTGT CCCAGGTGT CCCATGCTTC GTGCACGTT CCCATGCTTC CTCCAGGGG CCCATGCTTC CTCCAGGGG CCCATGCTTC CCCATGCTTC CCCCTTGGGGG CCCCTTGGCC CCCATGCTTC CCCCTTGGGGC CCCCTTGGGGC CCCCTTGTTCTC GTGCACGTTC CCCCTTGGGGC CCCCTTGTCTCT CTTCTTCTT	AACATGTATA GACCACAGCG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCACAAAG ACGAGTATTA GGAGGGCCGG TTGGTGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATA TTTGCTATTA TCTGTCCCAA GTTTGAATGT TCAGTCTCCCAA GTTTGAATGT TGATGTTTCC CAAGACACT CACTGTGGTC GGAATACACC CGACACTTT CACCTTTTT GGTCCGCATC ATGCTCATATC CAGGGGGTTT CACCTGTTTC CACTGTGTTC CACTGTGTT CTCCTATATC TCCTGTTGTGT TCCTGTTTC CCGAGCCTGT AGGTGATC CCGAGCCTGT CCGAGCCTGT CCCTATATC CCGAGCCTGT AGGTGTCCC CAAGGGGTCCC CGAGGCCTCT CCGAGCCTGT AGGTGTCCC CAAGGGGTCCC CGAGGCCTCC CAAGTGACACC CCGAGCCTTT CCCGAGCCTGT AGGTGACTCC CCAAGGGTCCC CGAGGCTCC CAAGTGACTCC CATCCATGTC CCTCCTGTTC CCGAGCCTCT CCGAGCCTCT CCGAGCCTCT CCGAGCCTCC CAAGTACTTC CCGAGCCTCC CAAGTACTTC CCAATGTCC CCAAGTCTCC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CACTCCATGTC CACCTCTTC CATCCATGTC CATCCATGTC CATCCATGTC CACCTCTTTT CACCTCTCTC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CATCCATGTC CACCTCTTC CATCCATGTC CATCCATGT CATCATT CATCCATGT CATCCATGT CATCCATGT CATCCATGT CATCCATGT CATCCATGT	120 180 240 300 360 420 600 600 780 840 900 1020 1140 1200 1140 1260 1320 1440 1500 1680 1740 1860 17920 1980 2040 2160 2220 2340 2460 2520
50 55 60 65 70	TCCAAAGTTC CACAGAAGTTC CACAGAAAGTTG GTGTAAAGAA CACTGCCTCC CCTTGGCAGA TTTTCTTGGC TCCTGGATGG GCTGCTCTT TCAGGCTGAC AAATGCCCTC TTTTTTTTTT	TTCCAGTCCT CTCAAATCAT AGCCAAATCAT AGCCAAATCAC CAGCAAAGGC TGGCCTTTAA TGTGGTGCAA TCCTCTCTCAGC TAGAGGCCAA TTCATTTCAC AATAAAAGAG TTAAATAAGAG TTAAACAAGC CTGGGGCCCC AGGGACCTCC CTGGGAGCAG ATTATGACCAA TCTTCTCCCC TTCTCACC TCTCAATGG AGAAGGTGTT ATGAGATGAT ATGAGATGAT ATGAGATGAT TCAACTTCCA TCTTCAAAA TCAAGACAGA ATTACAAAA TCAAGACAGA CCACGTTGGG ATTACAATAG ATCAAGACAGA ATTACAATAG ATCAAGACAGA ATCATTCTCTCCT GTATCAATAG ATCAAGACAGC CCGTCTTGCC CCGATTGGA ATCACACC CCGATTGTCA ATAACACC CCCCATTGTCC CCCAATCTCC CCCAATCTCC CCCAATCTCC CCCCAATCTCC CTTCTTCCC CTTCTTCCC CCCAATCTCC CTTCTTCCC CTTCTTCCC CTTCCAACC CCTTTGTCC CCTCCAACC CCTTTTTCCC CTTCTTTCCC CTTCTTTCCC CTTCTT	CCTAGGCATC AAGTGTACAG AGGACCGAA AGCACTATCC CATTTTTGTT GGCTGTGGAG GTGGAGGACT CCTTCATCCC AATATTCCCA ACTGGCCCTC GGCGACCAAC GTGTAACATT TGATCATAAA TCACAAGCCT TTTTGTGTTG TACAGAGTGC GACTTGATCAA AGTTGGCAAA AGTTGGCAAA AGTTGGCAAA CGAGCTTGGT CAACTTGGT CAACTTGGT CCACTTGGT GTTACACATT CTACAAGTGG GTACACATT CCTACAAGTGG GTACACATT CCTACTCCC GTCTGCTCCA CCCTTTCCT CGCCATCTGC GACCTTGGT GACCTTGTC GACCTTTCCT CGCCATCTGC GACCTTGCC GACCTTGCC GACCTTGCC GACCTTGCC GACCTTGCC GACCTTCCC CGCCACCTGC GACCTTCCC GGCCACCTGC GGCCACTGC GGCCACTGC GGCCACTGC GGCCACTGC GGCCACTGC GGCCACTGC GGCCAGCAG GGCCTGCTGT TAACTTGTTGTGC CAGCCAGCAG GGCCTGCTGT TAACTTGTAG	TTATTGATCC CTGATGAGTI TGTGAGCAGG GGACTTCTAA TAATTCAATT CTCAGGGTGG CCTGCCTTCC CTTCCCCTGC CAATTTTCTG TGCCTGCTC ACTAGGCAAA CTTTTAAAAT AGAGGGACAG AAACTTGTT AGGCAGAGG CAGCTCCTCT CTGCCAGAAG CCTCTCTCT CTGCCAGAAG CCTCTCTCTA ACAGAGTGAC AGCAGCATTTCA ACAGAGTGAT TCTTCCCTGG ACCAGAACT TCTTCCCTC GAAAATTTCT TCTTCCGTGA CCGGACTT TCCTCTCTCT CGGAAAATTTCT CCTCCGGACT TCCTCCTCC CGAAAATTTCT CCTCCGGACT CCTCAGACAG CCCCCTACCC CAGATTGTCA CCCCCTAGCC GAGTGTCCA CCCCTAGCC GAGTGTCCAGTGA GTCCCAGTGA GTCCCAGTGA GTCCCAGTGA GTCCCAGTGA GTCCCAGTGA GTCCCAGTGA GTCCCAGTGA	TCCAGTCGAG GTCAAAAAAT ACCTCAGTAG ACCATCGGT ATTCTTACTA ACTCCTGTTG CTGTTTAGAC CCCCAGTGCA GTCCTCTGG GTCATCCTGG CTAGGTCTTG TCACTCTGG TCACTCTGG CTAGGTCTTG TTCATTTTC CCTCTTATAA TTGGTTTGCT TTCATTTTC CCTCTTCCGAT GAGAGAAC CCTCTGGCAT GAGAGAAC CCTCTGGCAT TCGTTACAA GGATCCGGA TCTGTTACAA GGATCCGGA TCCTTGACAT TCGTTTCTC AGCTTGAAAT TCAGTTGCCG CCACGATGCT CCCCGCGTT CCCCCGTTC GCCCTCTCGCC CCCATGCTTC GCCCCTCTCGCC AGCGTTTTAAA GCCCCCCTTC GCCCCCCTTC CCCCCCTTC CCCCCCTTC CCCCCCTTCTCTCTC GCCCCCCTTC CCCCCCTTCTCTCTC	AACATGTATA GACCACAGG CCCCCTTTGT GAGTTTCATA ATCTTCTTCT GGCAGCCAGT ACCCACAAGA ACGAGTATTA GGAGAGGCCG TTGGTGGTGG GCTCAGACAT GTTTGTTGA CCCCAAAGAG TTTGCTATTA TCTGTCCCAA GTTTGATGTTGT CAAGTCACAT CCTGAACACT CAGACACT CAGACACT CAGACACT CACAGACACT CAGCACCTTT CACCTTTTT CACCTCTTT CACCTCTTT CACCTCTTT CACCTCTTT CACCTCTTT CACCTCTTT CAGGCGGTT CACCTATATC CCGAGCCTGT AAGTGATGGA TCAGGGGTCC CAAGAGTACTC CATCAGTGTC CATCATGTC CATCATGTC CATCATGTC CATCATGTC CATCATGTC CATCATGTC CATCATGTC CATCATGTC CTTCAATGTC	120 180 240 360 420 660 720 780 840 900 1020 1140 1260 1320 1380 1560 1680 1740 1880 1980 2040 2160 2220 2280 2340 2460

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	Coding seq	uence: 351.	.3701	~			
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Seq ID NO: 21 Protein sequence Protein Accession #: AAD04756

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55	AGTGCCTTGG	GGAATTTCCT	CTATTGATGT	ACAGTCTGTC	ATGAACATGT	TCCTGAATTC	4020
	TATTTGCTGG	GCTTTTTTTT	TCTCTTTCTC	TCCTTTCTTT	TTCTTCTTCC	CTCCCTATCT	4080
	AACCCTCCCA	TGGCACCTTC	AGACTTTGCT	TCCCATTGTG	GCTCCTATCT	GTGTTTTGAA	4140
		TGCCTTTAAA					4200
		ACTACTCTGT					4260
60							
00	GTTTAGAGAG	CTAAGATTAT	CIGGGGAAAI	CAAAACAAAA	AACAAGCAAA	CAAAAAAAAA	4320
	A						
	Seq ID NO:	25 Protein	sequence				
	Protein Acc	ession #: N	IP 000035.1				
65		*** -	_				
	1	11	21	31	41	51	
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	METIOT OF CO	VDDDDCvmvc	CARONE SOC.	PERTONDON	I DDDA A CA A CO	CASTITION	60
		YPRPPSKTYR					
70		QQQQQQQET					120
70		PEPGAAVAAS					180
	DILSEASTMQ	LLQQQQQEAV	SEGSSSGRAR	EASGAPTSSK	DNYLGGTSTI	SDNAKELCKA	240
	VSVSMGLGVE	ALEHLSPGEQ	LRGDCMYAPL	LGVPPAVRPT	PCAPLAECKG	SLLDDSAGKS	300
		KGGYTKGLEG					360
		GPPPPPPPPH					420
75		SSWHTLFTAE					480
, 5		GOESDFTAPD					540
		YYFPPQKTCL					600
		CPSCRLRKCY					660
00		PIFLNVLEAI					720
80	ALPGFRNLHV	DDQMAVIQYS	WMGLMVFAMG	WRSFTNVNSR	MLYPAPDLVP	NEYRMHKSRM	780
		SQEFGWLQIT					840
		SCSRRFYQLT					900
	VQVPKILSGK						5.0
	TOPMENTALD						

		26 DNA sequid Accession		uster			
5	TATCTCATCT	11   ATGGCCAGTG CAATCCTCTA	AATAACCATG	AAAGTTGATG	ATTATCTCAT	GGTACAGATG	60 120
10	NNTGAACCTG ATAATCTTGG CTCTCTCG CATAGACAGA	AGTGTTTAAT TGTTAATGGT ACTTTCAGAG CTCCTCTGTA GAGTAGTGAG	GTTTCTAGTC TATGAAGGAC ACAATTGGAG AAATATACTT	GATGCTGTTA GATTAAATAT AAACAGAGTT TTTTTAATAC	TCTGTTGCAC AACCCTTTGG CTAACAATAT AGAAGGTTCC	CACATTTTGA TATAAATGTT TAAAATCAGC CTGAAGTACT	180 240 300 360 420
15		AAAAAAAAAA		AATGAACAAT	TTTGGTCATA	AGCAGTTTCT	480
20	Nucleic Ac:	27 DNA sequid Accession tence: 64	#: NM_006	551.2			
20		11   GTCCAAATCA TGTCGGTGTG					60
25	AATGCCGAGT CCTCTGTTCA TTAGGAGTGA GTCCTGGTGA	TCTGCCCAGC AGTTAAGTCT AGAGATGCAC AAATATTGAA	TCTTGTTTCT TGCCAAATTT GGATCAGATG GAAATGTAGT	GAGCTGTTAG GATGCCCCTC TCCCTTCAGA GTGTGACATG	ACTTCTTCTT CGGAAGCTGT AACGAAGCCT TAAAAACTTT	CATTAGTGAA TGCAGCCAAG CATTGCGGAA CATCCTGGTT	120 180 240 300 360
30	GCTTTAATAA	TTCAATGACA ATCACTTGCT	CTAC	CACTGCAGAA	TGTAAAGGTT	TCAACGTCTT	420
35	Protein Acc	28 Protein cession #: 1	TP_006542.1				
33		11   TLALCCYQAN LQKRSLIAEV		31     LLDFFFISEP	41       LFKLSLAKFD	51   APPEAVAAKL	60
40	Nucleic Ac	29 DNA sequid Accession nence: 150	1 #: NM_0026	545.1			
45	1    -	11	21   CAGOGGATTE	31   ANACNATOTO	41   	51	60
45	ATGGCTCAGA ACAAGAGCAA AAACTGCAAA	 TATTTAGCAA AAGATGTGGA AGGATAGACA	 CAGCGGATTT CAAAGAAGAA AGTGACTGAC	 AAAGAATGTC GCATTACAGA AATCAGAGAG	 CATTTTCACA TGGAAGCAGA GCTTTGAGTT	 TCCGGAACCA GGCTTTAGCA GTCAAGCAGC	60 120 180
50	ATGGCTCAGA ACAGAGCAA AAACTGCAAA ACCAGAAAAA TCAGATTCCC GAGAAACTAT CCTATTCTGA	 TATTTAGCAA AAGATGTGGA AGGATAGACA AAGCACAGGT AAAAAAGAGC TGCTGGATGA GCCCTTCCTT	CAGCGGATTT CAAAGAAGAA AGTGACTGAC TTATAACAAG ATTAGATATT CAGTTTCGAG TTCAGCACAG	AAAGAATGTC GCATTACAGA AATCAGAGAG CAGGATTATG GATGTAGAAA ACTAAAAAAA CTCTATTTTA	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACCCA CACCTGTATT GACCTACTAT	TCCGGAACCA GGCTTTAGCA GTCTAGCAGC GTTTCCTGAA AGCTGAACTT ACCAGTTACT TCAGAGAGGA	120 180 240 300 360 420
	ATGGCTCAGA ACAAGAGCAA AAACTGCAAA ACCAGAAAAA TCAGATTCCC GAGAAACTAT CCTATTCTGA CAGTGGCCAC ACTTACAGTA TCTACAGAAC ACACCTGCCA	 TATTTAGCAA AAGATGTGGA AGGATAGACA AAGCACAGGT AAAAAAGAGC TGCTGGATGA	CAGCGGATTT CAAAGAAGAA AGTGACTGAC TTATAACAAG ATTAGATATT CAGTTTCGAG TTCAGCACAG TGGGCCTTCC ATTCCAAAAGT AAGTCTTCCG TCCACAAGGA	AAAGAATGTC GCATTACAGA AATCAGAGAG AATCAGAGAG CAGGATTATG GATGTAGAAA CTCTATTTTA ACTTATGCTT GGCTTCAATC GGACAATCTC AGCTTACCTA	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACCCA CACCTGTATT GACCTACTAT TACCTTCTAT CAAGAATGCC CATATTTCTC TCTATCGTCC	TCCGGAACCA GCCTTTAGCA GTCAAGCAGC GTTTCCTGAA AGCTGAACTT ACCAGTTACT TCAGAGAGGA TTATCCTTCT CACTTTTCCA ATATCCTTTG AGTAGTCAGT	120 180 240 300 360
50	ATGGCTCAGA ACAAGAGCAA AAACTGCAAA ACCAGAAAAA TCAGATTCCC GAGAAACTAT CCTATTCTGA CAGTGGCCAC ACTTACAGTAA TCTACAGAAC ACACCTGCCA ACTGACATGAAC ACACCTGCCA ACTGACATGG AAAGCAAGGA AAGCTAAGGG GTGGATAATG	TATTTAGCAA AAGATGTGGA AGGATAGACA AAGCACAGGT AAAAAAAGAGC TGCTGGATTACC AACAGGCTGC CTATATATTT CACCCTTTCA CAAAACTATT CTGAATTGGA ATATCAGTAA TGGAGGTAT TGGAGGTAT	CAGCGGATTT CAAAGAAGAA AGTGACTGAC TTATAACAAG ATTAGATATT CAGTTTCGAG TTCAGCACAG TGGGCCTTCC ATTCCAAAAT AAGTCTTCCG TCCACAAGGA TGACAAAATA GATAACAGAT ATTGACTGA AGACCTGAG	AAAGAATGTC GCATTACAGA AATCAGAGAG CAGGATTATG GATGTAGAAA CTCTATTTTA ACTTATGCTT GGCTTCAATC GGCAAATCTC AGCTTACCTA GCTAGCTACTT CCAAAAGTCA TCAAAAGTCA TTAGACTTGG GAAGAGAAAA	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACCCA CACCTGTATT TACCTTCTAT TACCTTCTAT CAAGAATGCC CATATTTCTC TCTATCGTCC CAGAATTTTT GCAATCTACA ATCCTCTAAG ATGTTCAAG	TCCGGAACCA GGCTTTAGCA GGCTTTAGCA GTCAAGCAGC GTTTCCTGAA AGCTGAACTT ACCAGTTACT TCAGAGAGGA TTATCCTTCT CACTTTTCCA AGTATCCTTTG AGTAGTCAGT AAAAATGGG GGTATCTCCA GTAGCCTAAG TTTGCTAGCA	120 180 240 300 360 420 480 540 600 660 720 780 840 900
50 55 60	ATGGCTCAGA ACAGAGACAA AACCTGCAAA ACCAGAAAAA TCAGATTCCC GAGAAACTAT CCTATTCTGA CAGTGGCCAC ACTTACAGTA ACACCTGCCA ACTGACACG AAGCAAGGA AAGCAAGGA AAGTCTGAGG GTGGATAATG AGGAACCTTACAGTAAGAAC ATTCGAACA ATTCGAACA ATTCGAACA	I TATTTAGCAA AAGATGTGGA AGGATAGACA AAGCACAGGT AAAAAAAGAGC GCCTTCCTT CTGGATTACC CTATATATTT CACCCTTTCA CAAAACTATT CTGATTTGGA ATATCAGTAA TGGAGGTATT GGGATGCATT CGGATGCAT CGGATGCAT CGGATGCT CTGATTTGGA ATATCAGTAA TGGAAAATC CTCAGCTTGC	CAGCGGATTT CAAAGAAGAA AGTGACTGAC TTATAACAAG ATTAGATATT CAGTTTCGAG TCAGCACAG TCAGCACAG TCACAAGGA TCACAAAGTA TTACAAAATA TATAACAAGT TCACAAAGTA TATAACAAGT TATAACAGAT TATAACAGAT TATTGACTGG AGACCATGAG TCTTCTTGAA ACTTTCTTGAA AAAAGCCCAG	AAAGAATGTC GCATTACAGA AATCAGAGAG CAGGATTATG GATGTAGAAA ACTAAAAAAA CTCTATTTTA ACTTATGCTT GGCTTCAATC GGACAATCTC AGCTTACCTA GCTTAGTACAT TCAAAAGTCA TTAGACTTG GAAGAGTCA TAGAGATCG GAAGAGTCA GAAGAGAAAA GAGAGAAAA GAGAGAATCGA GCAACTGTTA GCCATATATT	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACCCA CACCTGTATT TACCTTCTAT TACCTTCTAT CATGATCTC CATATTTCTC TCTATCGTC CAGAATTTTT GCAATCTACA ATCCTCTAAG ATGTTTCAAG ATGTTTCAAG CAGCAAATTG CAGCAAATTG CAGCAAATGA	TCCGGAACCA GCCTTTAGCA GCTCAAGCAGC GTTTCCTGAA AGCTGAACTT ACCAGTTACT TCAGAGAGGA TTATCCTTCT CACTTTTCCT AGTAGTCAGT AAAAATGGG GGTATCTCGAAAAAATGGG TTTGCTAGCA TCATCTTGAA TCATCTTGAA TCATCTTGAAG TCATCTTGAA TCATCTTGAAT CCCAAATGGG	120 180 240 300 360 420 480 660 720 840 900 900 1020 1080
50 55	ATGGCTCAGA ACAGAGACAA ACCAGAAAAA ACCAGAAACTATCCC GAGAAACTAT CCTATTCTGA ACACTGCAA ACTACAGAAC ACTACAGTAC ACACTGCCA ACTGCACA ACTGCACA ACTGCACAGA ACACTGCGAAAC ACACTGCAAAC ACACGTAGATC AGAAAGGTAGA AAGTAGATCTT AGAAAAGGTAGAAAC ACCAGTAGTT ATGGAACAA ACCAGTAGTT ATGGAAACC AATGCTAGTG	TATTTAGCAA AAGATGTGGA AGGATAGACA AAGCACAGGT AAAAAAAGAGC GCCTTCCTT CTGGATTACC AACAGGCTGC CTATATATTT CACCCTTTCA CAAAACTATT CTGATTTGGA ATATCAGTAA TGGAGGTATT GGGATGCTGC TCCAGCTTGC TCCAGCTTGC TCCAGCTTGC TCCAGCTTGC TCCAGCTTGC TCCAGCTTTT TGAAGGTCTT TGAAGGTCTT	CAGCGGATTT CAAAGAAGAA AGTGACTGAC TTATAACAAG ATTAGATATT CAGTTTCGAG TCAGCACAG TGGGCCTTCC ATTCCAAAAGT AGTCATCCGA TGACAAAGT TGACAAAGT ATTTGACTGG AGACCATGAG TCTTCTTGAC ACTTCTTGTG AAAAGCCCAG AAGTTCTCTT CATTACAAAGT CATTACAAAGT CATTACAAAG CATTACAAAG CATTACAAAG CATTACAAAGT CATTACAAAGT CATTACAAAGT CATTACAAATT CATTACAAAGT CATTACAAATT	AAAGAATGTC GCATTACAGA AATCAGAGAG AATCAGAGAG CAGGATTATG GATGAAAAAA CTCTATTTTA ACTTATGCTT GGCTTCAATT CAGAAATCCT AGCTTACCTA GCTAGTACAT TCAAAAGTCA TTAGACTTG GAAGAGTCA GAAGAGTCAA CAGAGAGATCAA CAGAGATCAT CAAAAGTCA TTGAAAACTCT CAAAAGTCA TTGAAGACT GCCATATTAT CTTCAAGAAG TTGAAGACCG GAAGGATTTC	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACCCA CACCTGTATT TACCTTCTAT TACCTTCTAT TACCTTCTAT TACCTTCTAT TCTATCGTC CAGAATTTTC CAGAATTTTT GCAATCTACA ATCTTCTAAG ATGTTCAAG ATGTTTCAAG ATGTTTCAAG CAGCAAATTG CAAGAAGCA CTCAGAAAGA TTGAAGTACA AATTTCCATA AATTTCCATA AATGAACAT AGCTACCAGT	TCCGGAACCA GCCTTTAGCA GCCTTTAGCA GCTCAAGCAGC GTTTCCTGAA AGCTGAACTT ACCAGTTACT TCAGAGAGGA TTATCCTTCT CACTTTTCCT AGTAGTCAGT AAAAATGGG GGTATCTCTAAAAAATGCAT TCCTTGAAA GCCTAAG TCATCTTGAA GTCTTTAAAT CCCAAATGGG GAATGAGGAG TACCAATCAC ATGCGGAGAA TACCTTTTACG	120 180 240 300 360 420 480 540 660 720 840 900 900 1020 1080 1140 1200 1260 1320
50 55 60	ATGGCTCAGA ACAGAGACAA ACCAGAAAAA ACCAGAAACTAC CGAGAAACTAT CCTATTCTGA CAGTGGCCAC ACTTACAGTAA TCTACAGGAAC ACACCTGCCA ACTGACAGGA ACACCTGCCA ACTGACAGGA AAGCCAGGA AAGTCTGAGG GTGGATAATG AAGGATCCTT AGAAAAGGTACTT ACGAACAA ACCAGTAGTT ATGGCAGCTT ATGGCAGCTT ATGGCAGCTT ATGGCAGCTT ATGCTAGTG TGTGAATGTG AGATCTGAATCC AATGCTAGTG TGTGAATGTG AGACTGAATAC CTGCAGAATA	I TATTTAGCAA AAGATGTGGA AGGATAGACA AAGCACAGGT AAAAAAGAGC GCCTTCCTT CTGGATTACC AACAGCTTCA CAACAGCTTCA CAACAGCTTCA CAACAGCTTCA CAACAGCTTCC CAACAGCTTCC CAACACTATT CGGATACATTT GGATGCTGC ATGCAACTGT ATGCAACTGG TTCTCACTTT CAGCTTTC CAGCTTTC CAGCTTTC CAGCTATT TGAAGGTCTC CAGCCTATTT TGAAGGTCTC CAGCCTATTT TGAAGGTCTC AAGTAGATGT AATTAGATGT AATTAGATGT ATCATTGCCT	CAGCGGATTT CAAAGAAGAA AGTGACTGAC TTATAACAAG ATTAGATATT CAGTTCGAG TCAGCACAG TGGGCCTTCC ATTCCAAAAATATA AGTCTTCCGA TGCACAAGGA TGACAAAATAA AATTTGACTGG AGACCATGAG TCTTCTTGAG AAAGCCCAG AAGTTCTTTTGTG AAAAGCCCAG AAGTTCTTTTCAAAA TTTACAAAA TTTACAAAT TTGGCAGCTAT	AAAGAATGTC GCATTACAGA AATCAGAGAG AATCAGAGAG CAGGATTATG GATGTAGAAAA ACTCAAATCT GGCTTCAATCT GGCTTACCTA GCTTAGCTT AGCTTACCTA GCTTAGCTT AGCTTACCTA GCTAGTACAT CTCAAAAGTCA TCAAAAGTCA TCAAAAGTCA GCAACTGTTA GCCATATAT CTTCAAGAAG TTGAAGACCA GTACAAGCGC GAAGGATTTC ATAATGCAAG GGTTCTAAAAG GAGCATATTC	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACTAT CACCTACTAT TACCTTCTAT TACCTTCTAT TACCTTCTAT TCATGGTC CAGAATTTTC CAGAATTTTCAC ATCTTCAAC ATCTTCAAC ATCTTCAAC ATCTTCAAC ATCTTCAAC ATCTTCAAC ATCTTCAAC ATTTCAAC CAGCAAATTC CAAGAAGAC CTCAGAAAGA TTGAAGTACA AATTTCCATA AAAGAAACAT AGCTACCAGT CCCTTTGCTG TTTTGTGGTCA AAAACTGTCC AAAACTGTCC AAAACTGTCC	TCCGGAACCA GCTTTAGCA GCTGAACTAC GTTACCTGAA AGCTGAACTAC TCAGAGAGGA TTATCCTTCT CACTTTTCCA ATATCCTTTC AGTAGTCAGT AAAAATGGG GGTATCTCA TCAGCCTAAG TCATCTTGA TCATCTTGA TCATCTTGA TCATCTTGA TCATCTTGA TCATCTTGAA TCATCTTTAAAT CCCAAATGGG GAATGAGGAGA TACCTTTTACG GGTACATTACG GGTACATTATA TAGAGGAACTG AAAATGGGAC	120 180 240 300 360 420 660 720 840 900 1020 1080 1140 1260 1320 1380 1440 1500
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	ATGGCTCAGA ACAGAGAGA AAACTGCAAA ACCAGAAAAA TCAGATTCCC GAGAAACTAT CCTATTCTGA CAGTGGCCAC ACTTACAGAAC ACTGACATGC AAAGCAAGGA AAGCAAGGA AAGCAAGGA AAGCAAGGA AAGCATCTGACAG AAGCATCTT AGAAAGGTGA ATTCGAACAA ACCAGTAGTT ATGGCAGATT TCGCACAAACC CATGCTAGTG GACTTGAATC GACTTGAATC GACACAATC CCCAGAATTA GCAGAAATTA GCAGAAAGTT GCAGAAAGTT GCAGAAAGTT GCAGAAAGTT GCAGAAAGTT	I TATTTAGCAA AAGATAGGACA AAGATAGACA AAGCACAGGT AAAAAAAGAGC TGCTGGATTACC AACAGGCTGC CTATATATTT CACCCTTTCA CAAAACTATT CTGATTTGGA ATATCAGTAA TGGAGGTATT GGGATGCTGT TGCAACTGG TTTGTCAGTTC CTCAACTGG TTTGTCGATTC GGCTACTGT TGAAGGTTCT TGAAGGTTCT TGAAGGTTCT TGAAGGTTTT TGATTACATTGC TTCTACTGT AAGTAGATCT ATCAATTGCA ATGAAACACC	CAGCGGATTT CAAAGAAGAA ATTAGATATT CAGTTTCGAG TTCAGCACAG TGGGCCTTCC ATTCCAAAAT AAGTCTTCCG TGCACAAAAT AAGTCTTCGAG TGACAAAATA ATTAGACAGA TGACAAAATA CATTACTGAG ACACATGAG TCTTCTTGAA CCTTTCTTGTA ACTTTCTTTTCATAAGCCCAG AAGTCTCTT CATTACAAAA GTTAAGTCCA CATTACATT TGGCAGCTTT TGGCAGCTTT TGGCAGCTTT CTTGACCTTC CTTGACCTTC CGTGGATTTA	AAAGAATGTC GCATTACAGA AATCAGAGAG CAGGATTATG GATGTAGAAA ACTAAAAAA ACTAATTTTA ACTTATGCTT GGCTTCAATC GGACAATCTC AGCTTACCTA GCTAGCTTACCTA GCTAGCTTACCTA GCTAGCTTACCTA GCTAGCTTAC GCAAATGTCA GCAAACTGTA GGCAAATGTA CTCAAGAGA TTGAAGACCA TTGAAGACCA TTGAAGACCA GTCACAGGGC GAAGAGTTC ATAATGCAAG GTTCTAAAAG GTTCTAAAAG GTTCTAAAAG GAGCATATTC ATAATGCAAG GTTCTAAAAG GAGCATATTC ATGAGCATGT AACAAACACC	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACCCA CACCTGTATT TACCTTCTAT TACCTTCTAT TACCTTCTAT TACCTTCTAT TACCTTCTAT TACCTTCTAT CAAGAATGCC CAGAATTTTT GCAATCTACA ATCTTCAAG ATGTTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATTTCCATA AAAGAAACAT CCAGAAAGA TTGAAGATACCA TTGAAGTACCA TTGTGTCC TTTGTGGTCA AAAACTGTCG GTCAAAATCT TGTAATCAAAT	TCCGGAACCA GGCTTTAGCA GGCTTTAGCA GTCTAGCAG GTTTCCTGAA AGCTGAACTT ACCAGTTACT TCAGAGAGGA TTATCCTTCT CACTTTTCCA ATATCCTTCT AAAAATGCG GGTATCTCCA TAAGCCTAAG TCTTTGAA GTCTTTAAAT CCCAAATGGG GAATGAGGAGA TACTTTTACA ATGCGGAGAA TACTTTTACG GGTACATCAT AGGGAATGAT AGAGGAAGTG	120 180 240 300 360 420 600 660 720 780 840 900 1020 1140 1200 1140 1320 1380 1440 1560 1560
50 55 60 65 70	ATGGCTCAGA ACAGAGACAA ACCAGAAAAA ACCAGAAACTATCCC GAGAAACTATCTGA ACTTACAGTAC ACTTACAGTAC ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAAACC AATGCAACAA ACCAGTAGTT ACGAACATAGT ATGGCAGCTT ATGGCAGCTT ATGGCAGCTT ATGCAGAACC AATGCTAGTG GATTACAGTAGTA CCAGAAATTA ACAGAAATTA ACAGAAATTA ACAGAAATTA ACAGAAATTA ACAGAAATTA ACAGAAATTA ACAGAAATTA ACAGAAATTA ACAGAAATTA GCAGAAGTAG GTAGAACCG	I TATTTAGCAA AAGATGTGGA AGGATAGACA AAGCACAGGT AAAAAAGAGGC GCCTTCCTT CTGGATTACC AACAGGCTGC CTATATATTT CACCCTTTCA CAGAACTATT CTGATTGGA ATATCAGTAA TGGAGGTATT GGGATGCTGC TCTCAGCTTGC TCCAGCTTGC TGCCAACTGG TTTGTCGATC CAGGCTATT TGAAGGTCTC GTTCTACTGT AAGTAGATGA ATCATTGCCT GATCAACTG GTCAACTG GTCAACTG CTCAACTG CTCAACTG CTCAACTG CTCAACTG CTCAACTG CCCATGAACCC CCATGAACCC CCATGAACCAC CCCATGACCAC CCCATGACCAC CCCTTCAAAT	CAGCGGATTT CAAAGAAGAA AGTGACTGAC TTATAACAAG ATTAGATATT CAGTTTCGAG TTCAGCACAG TGGGCCTTCC ATTCCAAAAGT AGTCTTCCAG TCACAAGGA TGACAAAGTA TATTAGACTGG AGACCATGAG TCTTCTTGAC AAAGCCCAG AAGTTCTTTT CATTACAAAG AAAGCCCAG AAGTTCTCTT CATTACAAAG CTTTACCAAA CTTTACCAAG CTTTACCAAG CTTTACCAAT TGGAAATCATT TGGCAGCTTT TGGAAGTCAT CGTGACTTT TGGACCTTC CGTGGATTTC CGTGGATTTC CGTGGATTTC CGTGGATTTC TGGAAAACCAA	AAAGAATGTC GCATTACAGA AATCAGAGAG AATCAGAGAG CAGGATTATG GATGTAGAAA ACTAAAAAAAA CTCTATTTTA ACTTATGCTT GGCTTCAATT CAGAAAGTCA GTAGAAATCA TTAAAAGTCA GTAGAAACTCA GCAACAGTGTA GCAACTGTTA GCCATATAT CTCAAGAGA CAGAGACAC GTCACAGCGC GAAGACCA GTTCTAAAA GTTCAAAAGTCA GTTCAAAAGTCA GCAATATT CTCAAGAGC GTACAAGCAC GTACAAGCAC GAAGAACTCT AACAAACAC CAAGAAACTCT CACCGAGCAG	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACTAT CACCTACTAT TACCTTCTAT TACCTTCTAT CAAGAATGCC CAGAATTTTT GCAATCTACA ATCTCTAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCATA AAGAAAGCA AATTTCCATA AAAGAAGCA AATTTCCATA AAAGAACAT AGCTACCAGT CCCTTTGCTG TTTGTGGTCA AAAACTGTCG GTCAAAATCT TGTAACAATCT TAGATCAAGT TAGATCATA TAGATCATA TAGATCAAGT	TCCGGAACCA GGCTTTAGCA GGCTGAACTA AGCTGAACTA AGCTGAACTA TCAGAGAGGA TTATCCTTCT CACTTTTCCT ACTATTCCTTCT AGTAGTCAGT AAAAATGGG GGTATCCCA TCAGCCTAAG TCATCTTGAA TCATCTTGAA TCATCTTGAA TCATCTTGAA TCATCAGCA TCAATCTTGAA TCCAATCAG GAATGAGGA TACCAATCAG TACCAATCAG GGTACATCAT AGAGGAAGT AGAGGAAGT AGAGGAAGT AGAAGAACCA AGAAAAACCT TCACCAACCAA AATTAAAGCT	120 180 240 300 360 420 660 720 840 900 1020 1080 1140 1260 1320 1380 1440 1500 1560 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	ATGGCTCAGA ACAGAGACAA ACAGAAAAA ACCAGAAAAA TCAGATTCCC GAGAAACTAT CCTATTCTGA CAGTGGCAC ACTTACAGTA TCTACAGAAC ACACAGAC ACACACAGG AAAGCAAGG GTGGATAATC AGGATCCTT AGAAAGGTGA ATTCGACAA ACCAGTAGTT AGAAAGGTGA ATCGACAAC CAGCACC AATGCTAGTG TGGATGAATC TGCAGAATA ACAGAACT CGCACAAAC CCAGAATA TGCAGAATA ACAGAAATTA ACAGAAATTA ACAGAAAGTG TGCAAAGATG TGCAAAGAATG GTACAAACTG GTACAAACTG GTACAAACTG GTACAAACTG GTACAAACTG GTACAAACTG GTACAAACTG GTACAAACTG	I TATTTAGCAA AAGATGTGGA AAGATAGACA AAGCACAGGT AAAAAAAGAGC GCCTTCCTT CTGGATTACC AACAGGCTGC CTATATATT CACCCTTTCA CAAAACTATT CTGATTGGA ATATCAGTAA TGGAGGTATT GGGATGATT GGGATGATT GGGATGTGC TCCAGCTTCC CAGCTTCC AGCACTGG TTTGTCGATC CAGGCTATT TGAAGGTCTC GTTCTACTGT AAGTAGATT ATCATTACC CAGGCTATT TGAAGGTCTC CAGGCTATT TGAAGGTCTC CAGGCTATT TGAAGGTCTC CAGGCTATT ATCATTACACT ATCATTACACT ATCATACACT ATCATACACT ATGAAACACC CCATGACCAA TCTTACACT CTCTACACT CTCTACACT CTCTACACT CTCTTACACT CTCTTACACT CTCTTACACT CTCTTACACT CTCTTACACT	CAGCGGATTT CAAAGAAGAA AGTGACTGAC TTATAACAAG ATTAGATATT CAGTTTCGAG TCAGCACAG TGGGCCTTCC ATTCCAAAATA AGTCATCCGA TGACAAAATA GATAACAAGA TCTTCTTGAC AGACCATGAG CCTTCTTGAC AAAGCCCAG AAGTTCTCTT CATTACAAAA CTTAAGTCCA CATTACAAAA TTTAGACTGT CATTACAAAA TTTAGACTCT CGTGAGTCCA CATTGACATT CGCAGCTAT TGGCAGCTAT TGGCAGCTAT CCTGGAGTCAT CCTGGAGTCAT CCTGGACTCTC CCTGGATTTA ACACCCTGTT TGAAAACCAA TTTAGATGGT	AAAGAATGTC GCATTACAGA AATCAGAGAG CAGGATTATG GATGTAGAAA CTCTATTTTA ACTTATGCTT GGCTTCAATC AGCTTACCTA GCTAGTACAT GCTAGTACAT GCTAGTACAT GCTAGTACAT GCAAAGTCC GAAGAGTCG GAAGAGTCGA GCACACTGTTA CTTCAAGAAG CTTCAAGAG GCACATATTT CTTCAAGAAG GTTCAACAGCG GAAGGATTTC ATAATGCAAG GTTCTAAAAG GTTCTAAAAG GTTCTAAAAG CTACAGCGC GAAGGATTTC ATAATGCAAG GTCCAATATT CTCAAAAGTC CTCCAAGAC CTCCAAGACTC CACAGACAC CTCCACAGCAC GAAGAACTC CACCAGACAC CTCCACAGCAC GTCGAGCAC GTCGAGACTC CCCAGAGCAC GTCGAGACTC GCAGGACTC GCAGGACTC GCAGGACTC GCAGGACTC GCAGGACAC GTCGAGACTC GCAGGACAC CCCAGGACAC CTCGAGACAC CCCAGGACAC CCCAGGAC CCCAGC CCCAGGAC CCCAGC CCCAGGAC CCCAGGAC CCCAGC CCCACAC CCCAGC CCCACAC CCCA	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACCA CACCTGTATT GACCTACTAT TACCTTCTAT TACCTTCTAT TACCTTCTAT TACCTTCTAT CAAGAATGCC CAGAATTTTT GCAATCTACA ATCTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATGAAAACCA AATTTCCATA AAGAAACCA TTGAAGAACCA TTGTGGTCA TTTGTGGTCA TTTGTGGTCA TTGTGTCG GTCAAAATCT TAGATCAAAT TAGATCAAAT TAGATCAAAT TAGATCAAAT TTGCATTA	TCCGGAACCA GGCTTTAGCA GGCTTTAGCA GTCTAGCAGCAGC GTTTCCTGAA AGCTGAACTT ACCAGTTACT TCAGAGAGGA TTATCCTTCT CACTTTTCCA ATATCCTTCT AAAAATGCG GGTATCCTAAA TTTGCTAGCA TCATCTTGAA GTCTTTAAAT CCCAAATCGC GAATGAGGAG TACCAATCAC ATGCGGAGAA TACTTTTACA ATGCGGAGAA TACTTTTACAT AGAGGAAGTG AAAATGGGAC GGCCCGAACA GGCCCGAACCT AGAAAAACCT TCACAACCAA	120 180 240 300 360 420 480 540 660 720 840 900 900 9140 1200 1320 1380 1440 1500 1560 1620 1620 1680
50 55 60 65 70	ATGGCTCAGA ACAGAGACAA ACCAGAAAAA ACCAGAAACTAT CCTATTCTGA CAGTGGCCAC ACTTACAGTA TCTACAGGAA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAA AAGATCTT AGAAAGGTAATG AGATCTT AGAAAGGTACAT ATGGCAGCTT ATGGCAGCTT ATGGCAGCTT ATGGCAGCTT ATGGATGTGATTGTA CCGCACAAACC AATGCTAGTG CTGCAGAATA ACAGAAATTA ACAGAAATTA TCGCAGCATT TCGCAGAATA ACAGAAATTA TCGCAGAATA ACAGAAAATA TTTGGAAGCTAA TTTGGAGGGAA TTTGGAGGAA	I TATTTAGCAA AAGATGTGGA AGGATAGACA AAGCACAGGT AAAAAAGAGC GCCTTCCTT CTGGATTACC ACACGCTTCC ATTCACTTC ATTCACTTA ATTCAGTAA TGGAGTATT TGAAGTCTC CTCACCTTC CTTCTACTGT ATTCACTT AAGTAGATGT ATCATTGCC GACTACAACT TCTACTTC CCCATGACAAC CCCATGACAAC CTCTTCAAAT TCTGTAGTGC CCATGACAAC CTCTTCAAAT TCTGTAGTGC AGAGACACTAC AAGACACTAC AAGACACTAC	CAGCGGATTT CAAAGAAGAA AGTGACTGAC TTATAACAAG ATTAGATATT CAGTTCGAG TTCAGCACAG TGGGCCTTCC ATTCCAAAAA AGTCATCTCGAG TGACAAAAT AAGTCTTCCGA TGACAAAGAT ATTTGACTGG AGACAATGAC ATTTGACTGG AAAGCCAGG AGTTCTTTGAG AAAGCCCAG AAGTTCTTTTCATAAAA ATTTACAAAA GTTAAGACCAT TGGCAGCTAT TGGCAGCTAT TGGCAGCTAT TGGCAGCTTT TGGAAATCAT TTGACCTTT TGAAACCAA TTTAGATCGT TTAGATCAT TTAGATCGT TTAGATCAT TTAGATCAAA TTTAGATCAT TTAGATCAT TTAGATCAT TTAGATCAT TTAGATCAC CAGGAGTTCA CAGGAGTTCA	AAAGAATGTC GCATTACAGA AATCAGAGAG AATCAGAGAG CAGGATTATG GATGTAGAAA ACTAAAAAAA CTCTATTTTA ACTTATGCTT GGCTTCAATC GGACAATCTC AGCTTACCTA GCTTACCTA GCTTACCTA GCTTACCTA GCTTACCTA GCTTACCTA GCTAGTACAT TCAAAAGTCA TTAGACTTGG GAAGAGAAAA CTAGAGACCA GTCACAAGCGC GAAGGATTTC ATAATGCAAG GTCCTAAAAG GAGCATATTC AGGCATATTC AGGAGATCT CACAGCGC GAAGAGTAAAA CACGGAGCTC CACCGAGCAC CTCCAGACAC CTCCAGACAC CTCCAGACAC CTCCAGACAC CTCCAGACAC CTCCAGACAC CTCCAGACAC CTCCAGACAC CTCGAGACTC CACCGAGCAC CTCGAGACTC AGGAGTAAAA ACTAGGGGCT	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACCAA ACCCTACTAT TACCTTCTAT TACCTTCTAT TACCTTCTAT TCATGGTC CAGAATTTTTC CAGAATTTTCAAG ATCTCTAAG ATCTCTAAG ATCTCTAAG ATCTCTAAG ATCTCTAAG ATCTCTAAG ATCTCTAAG ATCTCTAAG ATCTCTAAG ATTTTCAAG CAGCAAATTG CAAGAAGACA TTGAAGAGCA TTGAAGAGCA ATTTCCATA AAAGAACAT AGCTACCAGT TCTTTGGTCA AAAACTGTCG GTCAAAATCT TTGAATTCTTA TAGATTCATA TAGATTCATA TAGATTCATA TAGATTCATTA TAGATTCATTA TAGATTCATTA TAGATCAAGT TTGCCATTAC CACTTAATCC CACTTAATCC	TCCGGAACCA GGCTTTAGCA GGCTGAACTA AGCTGAACTA TCAGAGAGGA TTATCCTTCT CACTTTTCCA ATATCCTTTT CACTTTTCCA ATATCCTTTG AGTAGTCAGT AAAAATGGG GGTATCTCCA TCAGCCTAAG TCATCTTGAA GCCTAAG TCATCTTGAA GTCATCTTGAA GTCATCTTAAAT CCCAAATGGG GAATGAGGAGA TACTTTTACG GGTACATCAC AGAAGAGGAGT CACCAACCAA AGAAGAACCT TCACCAACCAA AATTAAAGCT AGAACCAA AATTAAAGCT AGAATCACTA AGATCACCT TCACCAA AATTAAAGCT AGAATCACTA AGATTAATCT TGAAAATCCT TGAAAATCCT	120 180 240 300 360 420 480 600 660 720 780 840 900 1020 1120 1260 1320 1320 1440 1500 1620 1680 1740 1800 1740 1800 1800 1900 1
50 55 60 65 70	ATGGCTCAGA ACAGAGACAA ACAGAAAAA ACCAGAAAAA TCAGATTCCC GAGAAACTAT CCTATTCTGA CAGTGGCAC ACTTACAGTA ACACCTGCCA ACTGACACA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAA ACACTGCAAACC AATGCAACA ACACAGAACT ATGGCAGAT ATGGCAGAT ATGGCAGAT GCACAAACC AATGCTAGTG GACTGAATA CCTGCAGAATA ACAGAAATTA ACAGAAATTA GCAGAAATTA ACAGAAATTA GCAGAAATTA ACAGAAATTA ACAGAAATTA ACAGAAATTA ACAGAAATTA ACAGAAATTA TTGGAAGAG GTAAAAAAAA AAAGACTAA ATTTGGAGGAG GTTCAAGTAA	I TATTTAGCAA AAGATGTGGA AAGATAGACA AAGCACAGGT AAAAAAAGAGC GCCTTCCTT CTGGATTAC CTGGATTAC CTGATTTC CAACAGCTTG ATATATTT CTGATTTGA ATATCAGTAA TGGAGGTATT TGGAGGTATT TGGAGTATT TGCAGTTGC TTCTACTGT TTGTCGATT TGAAGCTGT TTTGTCGAT TGAAGGTTT TGAAGGTAT TGAAGGTAT TGAAGTTC CAGGCTATT TGAAGGTAT TGAAGGTAT TCATGAAC TCAGCTTC CATCAACT TTCTACAGT ATGAAACC CTCATGACAC CCCATGACAGA CTCTTCAACT ATGAAACCC CCATGACAGA CTCTTCAACT AGGAGACACT AGAGAGCACT GACTACACT GAGAGACCAG GCATAAACCA	CAGCGGATTT CAAAGAAGAA AGTGACTGAC TTATAACAAG ATTAGATATT CAGTTTCGAG TCAGCACAG TGGGCCTTCC ATTCCAAAAGT AGTCATCCGA TCACAAGGA TGACAAAGTA TATTAGATGG AGACCATGAG TCTTCTTGAC AAAACCCAG AAGTCCTTCTTT TGGAAGTCAT TGGAAGTCAT TGGAAGTCAT CATTACAAAA CTTGACCTTT TGGAAGTCAT CATTGACTTT TGGAAGTCAT CTTGACCTTT TGGAAGTCAT CTTGACCTTT TGGAAGTCAT TTGACCTTT TGAAAACCAA TTTAGATGGT TAAATCTTCCA ATTAACTGCA ATTAACTGCA	AAAGAATGTC GCATTACAGA AATCAGAGAG CAGGATTATG GATGTAGAAA ACTAAAAAAA CTCTATTTTA ACTTATGCTT GGCTTCAATC GGACTACATA GCTAGTACATA GCTTACCTA GCTTACCTA GCTTACCTA GCTAGTACAT CTAAAAGTCA GTAGAGACTCA GCAATCTGG GAAGAGTCAA GCAACTGTA GCCATATAT CTTCAAGAAG GTACAAGCGC GAAGGATTTC ATAATGCAAG GTTCTAAAAG GTTCTAAAAG GTTCTAAAAG CTCCCAGCGC CAAGGATTTC AGGACACTCT CACCGAGCAC GAGGACTCT CACCGAGCAG GTCGAGACTCT CACCGAGCAG GTCGAGACTCT CACCGAGCAG GTCGAGACTCT CACCGAGCAG GTCGAGACTCT CACCGAGCAG GTCGAGACTCT GCCAATTTATG GCCAATTTATG GCCAATTTATG	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACCTA CACCTACTAT CACGTACTAT TACCTTCTAT CATGATCTC TCTATCGTC CAGAATTTTC CAGAATCTAC ATCTCTAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG CAGCAAATTG CAAGAAACAT AGCTACCAGT CCCTTTGCTG TTTGTGGTCA AAAACTGTCG GTCAAAAACT TAGATCTAAG TTAGATCAAGT TAGATCAAGT TAGATCA	TCCGGAACCA GGCTTTAGCA GGCTTTAGCA GTCTAGCAGCAGC GTTTCCTGAA AGCTGAACTT ACCAGTTACT TCAGAGAGGA TTATCCTTCT CACTTTTCCA ATATCCTTCT AAAAATGGG GGTATCTCCA TAGCCTAAG TTTGCTAGCA TCATCTTGAA GTCTTTAAAT CCCAAATGGG GAATGAGGAGA TACCAATCAC AGGGAGAA TACCTTTTACG GGTACATCAT AGAGGAACA AGGAAAACCT TCACAACCAA AATTAAAGCT TCACAACCAA GAAATCAGTA GAACTCTTTG	120 180 240 300 360 420 660 720 840 900 1020 1080 1140 1260 1320 1380 1440 1500 1680 1740 1880 1800 1980
50 55 60 65 70	ATGGCTCAGA ACAGAGACAA ACAGAAAAA TCAGATTCCC GAGAAACTAT CCTATTCTGA CAGTGGCCAC ACTTACAGATA TCTACAGATA ACTGACATGG AAAGCAAGGA AATTCTGAACAA ACCAGTAGTT TGTACAGATA TCTGCAGAATT ACCAGAATTA ACAGAAATTA ACAGAAATTA ACAGAAATTA TCTGCAGAATA TTTGGAGAGA TTTCGGAGATA ATTCTGGAGATA AATTCTGGTA AAGAAATTA AAGAAATTA ACAGAAATTA TTTGGAGGAG GTTACAACAG GTTACAACAG GTTACAACAG GTTACAACAG AATTCTGGTA AACTTCAGTA AATTCTGGTA AACTTCAGACAA	I TATTTAGCAA AAGATGTGGA AAGATAGACA AAGACACAGGT AAAAAAAGAGC TGCTGGATTAC CTGGATTACC AACAGGCTGC CTATATATTT CTGATTTGGA ATATCAGTAA TGGAGGTATA TGGAGGTATT GGGATGCTGT TCTGGATTTGGA TTGGATTTGGA TTGGAGTTTC CTCAGCTTGC TTCACTGT TGCAACTGG TTTGTCGATC GATTACATTA AGTAGGTATT TGAAGGTCTT AAGTAGATGC TTGACTGG TTCTACTGT AACACAC CCATGACAC CCATGACAC CCCATGACAC CCCATGACAC CCCATGACAC CCCATGACAC CCCATGACAC CCATGACAC CCCATGACAC AGGAGGCAC AGGAGGCAC AGGAGGCAC AGGAGCCTCA AGCAGCTCCA AGCAGCTCCCA	CAGCGGATTT CAAAGAAGAA ATTAGATATT CAGTTTCGAG TTCAGCACAG TGGGCCTTCC ATTCCAAAAT AAGTCTTCCG TGCACAAAAT AAGTCTTCCG ATTACAAAAT AATTACAAAAT AATTACAAAAT GATAACAGAT TCTTCTGTG AAAACCCAG AAGTCTCTT CATTACAAAA GTTAACTACT CGGAGCTAT TGGCAGCTAT TGGCAGCTAT TGGAAATCATT TGGCAGCTAT TGGAAAACCAA ATTAGATGAC TTTAGATGCT CATGACTTC CATGACTTC CAGGAGTTCA TTAAACTCCA AATTACTCCA AAGTCTCCA CAGGAGTTCA ATTAACTGCA ATTAACTGCA ATTAACTGCC CAGGAGTTCA TTTAACTGCC CTTTACTATT	AAAGAATGTC GCATTACAGA AATCAGAGAG CAGGATTATG GATGTAGAAA ACTAAAAAA ACTAATTTTA ACTTATGCTT GGCTTCAATC GGACAATCTC AGCTTACCTA GCTAGCTTACCTA GCTAGCTTACCTA GCTAGCTTACCTA GCTAGCTTAC TCAAAAGTCA TTAGACTTGG GAAGAGAAAA GAGAATCTTA GCCATATAT CTTCAAGAAG TTGAAGACCA GTCACAGCGC GAAGGATTAC GAGCATATTC CACCAGCAG GTCACAGCGC GAAGGATTCT AACAACACC GAAGAACTCT AACAAACAC GTCGAGACT AACAACACC GAAGAACTCT CACCGAGCAG GTCGAGACT ACCAGAGCAC GTCGAGACT ACCAGAGCAC GTCGAGACT CACCGAGCAG GTCGAGACT CACCGAGCAC TTGCTGCT CAAGTACC CAAGTACC TTGCTGCT CTCTCTCTC CTCTCTCT CTCTCTCTC	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACCA CACCTGTATT GACCTACTAT TACCTTCTAT CAAGAATGCC CATATTTCTC CAGAATTTTT GCAATCTACA ATCTCTCAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCAAG ATGTTCATA AAAGAAACAT CCAGAAAGA TCACAGA CCCTTTGCTG TTTGTGGTCA AAAACTGTCG GTCAAAATCT TGTATCAAAT TAGATCAAAT TAGATCAAAT TAGATCATA TAGATCATA CTGCCATTAC CTGCTGATGC CTGCTGATGC CTGCTGATGC CTGCTGATGC CACTTAATCC ATCTCTCAAA	TCCGGAACCA GGCTTTAGCA GGCTTTAGCA GGCTGAACTT ACCAGTTACT TCAGAGAGGA TTATCCTTCT CACTTTTCCA ATATCCTTCT CACTTTTCCA ATATCCTTCT AAAAATGGG GGTATCTCCA TAGCCTAAG TTTGCTAGCA TCATCTTGAA GCCTAAG TCATCTTGAA GTCTTTAAAT CCCAAATGGG GAATGAGGAGA TACCAATCAC GGCACAATCAC GGCCGAACA AGGAACATCA GGCCCGAACA AGGAACATCAC GGCCCGAACA AGGAACATCAC AATTAAAGCT TCACAACCAA AATTAAAGCT AGAACATGG AAGTAATTCG AAGGAAGCATGG AAGTAATTCG	120 180 240 300 360 420 480 660 660 780 840 900 1020 1020 1140 1260 1380 1440 1500 1680 1740 1680 1740 1860 1920 1980 2040
50 55 60 65 70	ATGGCTCAGA ACAGAGACAA ACCAGAAAAA ACCAGAAACTAT CCTATTCTGA CAGTGGCCA ACTGCACA ACACTGCCA ACTGACACA ACACTGCCA ACTGACAGAA ACACTGCAA ACACTGCCA ACTGACAGGA AAGCCAGGA AAGCAAGGA AATTCTGAG ATTCGAACA ACTGACAGA ATTCGAACA ACCAGTAGTT ATGGCAGCTT ATGGCAGCTT ATGGCAGCT CGCACAAACC AATGCTAGTG TGTAATGTGA ACCAGAAGTA CCTGCAGAATA ACAGAAATTA GCAGAAGTC GTAAGAAAAA ACTACAACAG GTTCAAGTA AATTCTGGTA ATTTCGAGGAG GTTCAAGTA AATTCTGGTA ACTACAACAG GTACAACTA GTACAACAG GTATCAACTAG	I TATTTAGCAA AAGATGTGGA AGGATAGACA AAGCACAGGT AAAAAAGAGC GCCTTCCTT CTGGATTACC AACAGCTTCA CTGATTACT CACACTTCA CAAAACTATT CACCTTTCA CAAAACTATT CTGATTTGGA ATATCAGTAA TGGAGGTATT GGGATGCTG CTCAGCTTGC TTCACTTT CAGCTTGC TTCACTTT TGAAGGTCTC CTGACTTGC GTTCTACTGT ATGAAACAC CCATGACAGC CTCTTCAAAT TCTGTAGTGC GCATACACC GCAGCACTAG CTCTTCAAAT TCTGTAGTGC GCATACACC CCATGACAGC CTCTTCAAAT TCTGTAGTGC GGAGACACTAG GCATAAACCA AGCAGCTCCA AGCAGCTCCA AGCAGCTCCA ATGAAAAAAAAAA	CAGCGGATTT CAAAGAAGAA AGTGACTGAC TTATAACAAG ATTAGATATT CAGTTCGAG TGGGCCTTCC ATTCCAAAGA TGACACAGG TGCACAAGGA TGACAAAGAA TATTAGATGG AGACCATGAG ACTTCTTGAG AAAGCCCAG AGGTCCTTCTTGAG AAAGTCTTCTTTTCAAAA GTTAACACAT CATTACAAAA GTTAACACT CATTGACTT TGGAAGTCAT TGGAAGTCAT TGGAAGTCAT TGGAAGTCAT TGGAAGTCAT TGGAAGTCAT TGGAAGTCAT TGAAAACCAA TTTAGATGGT TAATCTTCA ATTACAAAA TTTAGATGGT CAGGAGTTCA ATTAACTGCA AGACTGTTCC CATTGACTTC AGAAGTCTAC CAGGAGTTCA ATTAACTGCA AGACTGTTCC GTTTACTATT CTACTTGATA	AAAGAATGTC GCATTACAGA AATCAGAGAG AATCAGAGAG CAGGATTATG GATGTAGAAA ACTAAAAAAA CTCTATTTTA ACTTATGCTT GGCTTCAATC GGACAATCTC AGCTTACCTA GCTTACCTA GCTTACCTA GCTTACCTA GCTAGTACAT TCAAAAGTCG GAAGAGCATGT GGCAATCTG GCAACTGTTA GCCATATAT CTTCAAGAAG TTGAAGACCA GTACAACACCC GAAGGATTTC AGTGCAATGT AACAACACC GAAGAACTCT CACCGAGCAG GTCCAAAACTCC CACCGAGCAG GTCGAGACTC CACCGAGCAG GTCGAGACTC CACCGAGCAG TCGAGACTC CACCGAGCAG TCGAGACTC CACCGAGCAG TCGAGACTC CACCGAGCAG TCGAGACTC CACCGAGCAG TCGAGACTC CACCGAGCAG TTTGCATGCT CACAGCTC CACTATTATG CAAAGTACC TTTGCTCC TTTCACTGT TTTCACTGT TTTCACTGT	CATTTTCACA TGGAAGCAGA GCTTTGAGTT ATCTCATGGT AGCTCACTAT CACCTACTAT TACCTTCTAT TACCTTCTAT TACCTTCTAT TACCTTCTAT TACCTTCTAT TACCTTCAT TACCTTCTAT CAAGAATGCC CAGAATTTTTC CAGAATTTTCAG ATGTTCAGA ATGTTCAGA ATGTTCAGA ATGTTCAGA ATTTCCATA AAAGAAGCCA CTCAGAAAGA TTGAAGTACA AATTTCCATA AAAGAACCT TTTGTGGTC GTCAAAATCT TTGTACAGT TTGTGTGTG TTGTACAAGT TTGAGTCATACA TTGAGTTCTA TAGATTCACA TTGCCATTAC CTGCTGATGT CACTTAATCC ATCTTCTCAG AGAGTTCAA AATGTCAAATGT CACTTAATCC ATCTTCTCAG AGAGTTCACAATTGC CTCACAATGG	TCCGGAACCA GCCTTTAGCA GCTCTAGCA GCTGAACTT ACCAGTTACT ACCAGTTACT TCAGAGAGGA TTATCCTTCT CACTTTTCCA AGATACCTTC AGTAGTCAGT AAAAATGGG GGTATCTCTA AGAACTAAA TCTTTAAAT CCCAAATGGG GAATGAGGA TACCATTTACA AGCAATCAC AGGAGAATGAC AGGAGAATGAC AGAACAATGAC AGAACAACCA AGAAAACCT TCACAACCAA AGAAAACCT CACAACCAA AGAAAACCT CACAACCAA AGAAAACCT CACAACCAA AGAAAACCT CACAACCAA AGAAAACCT CACAACCAA AGAAAACCT CACAACCAA AGAAACCT CACAACCAA AGAAAACCT CACAACCAA AGAAACCT CACAACCAA AGAAACCT CACAACCAA AGAAACCT CACAACCAA AGAAACCT CACCACCA	120 180 240 300 360 420 480 600 660 720 780 840 900 1020 1120 1260 1320 1320 1440 1500 1620 1680 1740 1800 1980 2040 2040 2160 2160 2160

	CACCTTACTC	TTTTTGGAAT	TTTAAATCAG	AGCAGTGGAA	GTTCCCCTGA	TTCTAATAAG	2340
		GACCAGAAGC					2400
		GTGGAACTAA					
							2460
~		TTACCAAAAA					2520
5	CCTTCTCCTG	CATTTGATAT	TATTTATACA	ACTCCTCAAG	TTGACAGAAG	CATTATACAG	2580
	CAACATAACT	TAGAAACACT	AGAGAATGAT	ATAAAAGGGA	AACTTCTTGA	TATTCTTCAT	2640
		CACTTGGACT					2700
		AACACCCAAA					2760
10		TTGCCAAAAC					2820
10	ATTGCATTGG	AACTTCTTGA	TTCAAAATTT	GCTGATCAGG	AAGTAAGATC	CCTAGCTGTG	2880
	ACCTGGATTG	AGGCCATTAG	TGATGATGAG	CTAACAGATC	TTCTTCCACA	GTTTGTACAA	2940
		ATGAAATTTA					3000
		TCCAGATAGC					3060
1 ~	GTACAGTTTA	GTACCCGATA	CGAACATGTT	TTGGGTGCTC	TCCTGTCAGT	AGGAGGAAAA	3120
15	CGACTTAGAG	AAGAACTTCT	AAAACAGACG	AAACTTGTAC	AGCTTTTAGG	AGGAGTAGCA	3180
		GGCAGGCTAG					3240
		CCTTTTTTCA					
							3300
		TAAATATTAA					3360
	GTCACAATGG	TGAATGCTGA	CCCTCTGGGA	GAAGAAATTA	ATGTCATGTT	TAAGGTTGGT	3420
20	GAAGATCTTC	GGCAAGATAT	GTTAGCTTTA	CAGATGATAA	AGATTATEGA	TANGATOTCG	3480
		GACTAGATCT					3540
		TGGAGCTGGT					3600
	GGTGTGACAG	GATCCTTTAA	AGATAAACCA	CTTGCAGAGT	GGCTAAGGAA	ATACAATCCC	3660
	TCTGAAGAAG	AATATGAAAA	GGCTTCAGAG	AACTTTATCT	ATTCCTGTGC	TGGATGCTGT	3720
25		ATGTTTTAGG					3780
23							
		TGTTTCACAT					3840
	AGCTTCAAAA	GGGATCGGGC	TCCTTTTGTG	CTGACCTCTG	ATATGGCATA	TGTCATTAAT	3900
	GGGGGTGAAA	AGCCCACCAT	TCGTTTTCAG	TTGTTTGTGG	ACCTCTGCTG	TCAGGCCTAC	3960
		GAAAGCAGAC					4020
30							
50		AACTTACAAG					4080
	CAAACTACAG	ACGCAGAAGC	TACAATTTTC	TTTACTAGGC	TTATTGAATC	AAGTTTGGGA	4140
	AGCATTGCCA	CAAAGTTTAA	CTTCTTCATT	CACAACCTTG	CTCAGCTTCG	TTTTTCTGGT	4200
		ATGATGAGCC					4260
		TCAAGGAAGT					4320
35							
33		ATGTAGTCCG					4380
	CGAACATTTG	TCGAATTTCA	GGAACTTCAC	AATAAGCTCA	GTATTATTTT	TCCACTTTGG	4440
	AAGTTACCAG	GCTTTCCTAA	TAGGATGGTT	CTAGGAAGAA	CACACATAAA	AGATGTAGCA	4500
		AAATTGAGTT					4560
40		GTGATCTTGT					4620
40		CTAGGTCTGC					4680
	GGAGCTGTGA	AATTATCCAT	CTCTTACCGA	AATGGTACTC	TTTTCATCAT	GGTGATGCAT	4740
	ATCABAGATC	TTGTTACTGA	AGATGGAGCT	GACCCAAATC	CATATGTCAA	AACATACCTA	4800
		ACCACAAAAC					
							4860
15		ATGAAATGCT					4920
45	GAACTTCAAC	TAAGTGTACT	CAGTGCAGAA	TCTCTGCGGG	AGAATTTTTT	CTTGGGTGGA	4980
	GTAACCCTGC	CTTTGAAAGA	TTTCAACTTG	AGCAAAGAGA	CGGTTAAATG	GTATCAGCTG	5040
		CATACTTGTA					
			••				
	0 TD 170	30 Duch-					
50		30 Protein					
50		30 Protein cession #: N					
50							
50	Protein Acc	ession #: N	IP_002636.1	31	41	51	
50				31	41	51	
50	Protein Acc	cession #: N 11 	7P_002636.1 21 	]	1	1	
	Protein Acc	cession #: N 11     KECPFSHPEP	IP_002636.1 21   TRAKDVDKEE	] ALQMEAEALA	   KLQKDRQVTD	) NQRGFELSSS	60
50 55	Protein Acc	cession #: N 11 	IP_002636.1 21   TRAKDVDKEE	] ALQMEAEALA	   KLQKDRQVTD	) NQRGFELSSS	60 120
	Protein Acc	cession #: N 11     KECPFSHPEP   QDYDLMVFPE	IP_002636.1  21  TRAKDVDKEE SDSQKRALDI	ALQMEAEALA DVEKLTQAEL	 KLQKDRQVTD EKLLLDDSFE	) NORGFELSSS TKKTPVLPVT	120
	Protein Acc 1     MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ	cession #: N 11     KECPFSHPEP   QDYDLMVFPE   LYFRPTIQRG	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS	 ALQMEAEALA DVEKLTQAEL TYALPSIYPS	 KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN	) NQRGFELSSS TKKTPVLPVT GFNPRMPTFP	120 180
	Protein Acc	11     KECPFSHPEP  QDYDLMVFPE  LYFRPTIQRG  GQSPYFSYPL	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS	 KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI	) NQRGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG	120 180 240
	Protein Acc	11	IP_002636.1  21  TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK	 KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA	120 180 240 300
55	Protein Acc 1   MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTOLEITD KOPWDAVLLE	11 	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN	 KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ	) NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG	120 180 240
	Protein Acc 1   MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTOLEITD KOPWDAVLLE	11	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN	 KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ	) NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG	120 180 240 300
55	Protein Acc 1   MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KOPWDAVLLE TSSLPTGSSL	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE	P_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH	KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP	) NQRGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE	120 180 240 300 360 420
55	Protein Acc 1   MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KOPWDAVLLE TSSLPTGSSL NASVKVSIDI	11   	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSX MAAFCRSITK CDVSTVEII	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD	KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY	) NQRGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV	120 180 240 300 360 420 480
55	Protein Acc 1   MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KOPWDAVLLE TSSLPTGSSI NASVKVSIDI LQNNHCLGSH	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWFPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITE CDVSSTVEII TEIRLQLLTF	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYINH LMQALCWVHD SAMCQNLART	KLQKDRQVTD KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP	120 180 240 300 360 420 480 540
55	Protein Acc 1   MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KDPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNNHCLGSH CKEAMTRHPV	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD EELLDSYHNQ	P_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KKSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH LKTKFPYTNH SAMCQNLART HRAVDQVIKA	KLQKDRQVTD KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV	120 180 240 300 360 420 480
55 60	Protein Acc 1   MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KDPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNNHCLGSH CKEAMTRHPV	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD	P_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KKSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH LKTKFPYTNH SAMCQNLART HRAVDQVIKA	KLQKDRQVTD KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV	120 180 240 300 360 420 480 540
55 60	Protein Acc 1   MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KOPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNNHCLGSH CKEAMTRHPV KKLKRAVNLP	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD RSKTADVTSL	IP_002636.1  21  TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW KKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF TEIRLQLLTF VELALQIENQ PGGEDTSRSS	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART HRAVDQVIKA TRGSLNPENP	KLQKDRQVTD KLQKDRQVTD KLQLDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVUD VRKICSALDG VQVSINQLTA	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA	120 180 240 300 360 420 480 540 600
55	Protein Acc 1   MAQIFSNSGF TERKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KOPWDAVLLE TSSLPTGSSL TASVKVSIDI LQNNHCLGSH CKEAMTRHPV KKLKRAVNLP NSGRSPTDCA	11	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGDS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENF PGGEDTSRSS TTTEQLOFTI	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART HRAVDQVIKA TTGSLINPENP FAAHGISSNW	KLQKDRQVTD KLQKDRQVTD KLQLDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSILLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESP VETLAITESP CSLSHNGKDL	120 180 240 300 360 420 480 540 600 660 720
55 60	Protein Acc 1   MAQIFSNSGF TERKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KOPWDAVLLE TSSLPTGSSL LASVKVSIDI LQNNHCLGSH CKEAMTRHPV KILKRAVNLD PNSGRSPTDCA FKPIQSKKVG	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE EGFQLPVTFT EHIQNCRKWD EELLDSYHNQ RSKTADVTSQ CSSKSVKEAW TYKNFFYLIK	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQFTI WDELIIFPIQ	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH LMQALCWVHD SAMCQNLART HRAVDQVIKA TRGSLNPENP FAAHGISSNW ISQLPLESVL	KLQKDRQVTD KLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VIKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSFDSNK	120 180 240 300 360 420 480 540 600 720 780
55 60	Protein Acc  1	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD EELLDSYHNQ RSKTADVTSL QSSKSVKEAW TYKNFFYLIK VSLPLCDFRR	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQPTI WDELIIFPIQ PLTCGTKLLY	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH LMQALCWVHD SAMCQNLART HRAVDQVIKA TRGSLNPENP FASHGISSNW ISQLPLESVL LWTSSHTNSV	KLQKDRQVTD KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ PGTVTKKGYV	NQRGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRHA CSLSHNKKDL SSGSSPDSNK MERIVLQVDF	120 180 240 300 360 420 480 540 600 720 780 840
55 60	Protein Acc 1   MAQIFSNSGF TERKAQVYNIA PILSPSFSAQ STEPIYLSLP KARTDLEITD KDPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNNHCLGSH CKEAMTRHPV KKLKRAVNLP NSGRSPTDCA FKPIQSKKVG QRKGPEALGK PSPAFDIIYT	11    KECPFSHPEP   QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD EELLDSYHNQ RSKTADVTSL QSSKSVKEAW TYKNFFYLIK VSLPLCDFRR TPQVDRSIIQ	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF TEIRLQLLTF PEGEDTSRSS TTTEQLQFTI WDELLIFPIQ WDELLIFPIQ PLTCGTKLLY QHNLETLEND	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLDLILH	KLQKDRQVTD KLQKDRQVTD KLQLDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ RGTVTKKGYV RDSSLGLSKE	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSILLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWEKRY	120 180 240 300 360 420 480 540 600 720 780
<ul><li>55</li><li>60</li><li>65</li></ul>	Protein Acc 1   MAQIFSNSGF TERKAQVYNIA PILSPSFSAQ STEPIYLSLP KARTDLEITD KDPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNNHCLGSH CKEAMTRHPV KKLKRAVNLP NSGRSPTDCA FKPIQSKKVG QRKGPEALGK PSPAFDIIYT	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD EELLDSYHNQ RSKTADVTSL QSSKSVKEAW TYKNFFYLIK VSLPLCDFRR	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF TEIRLQLLTF PEGEDTSRSS TTTEQLQFTI WDELLIFPIQ WDELLIFPIQ PLTCGTKLLY QHNLETLEND	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLDLILH	KLQKDRQVTD KLQKDRQVTD KLQLDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ RGTVTKKGYV RDSSLGLSKE	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSILLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWEKRY	120 180 240 300 360 420 480 540 600 720 780 840
<ul><li>55</li><li>60</li><li>65</li></ul>	Protein Acc  1	11    KECPFSHPEP   QDYDLMVFPE   LYFRPTIQRG   GQSPYFSYPL   SKVSNLQVSP   ERSTANCHLE   LQEVEVQNEE   LQEVEVQNEE   LGFQLPVTFT   EHIQNCRKWD   EELLDSYHNQ   RSKTADVTSL   QSSKSVKEAW   TYKNFFYLIK   VSLPLCDFR   KILASAPNWK	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIEND PGGEDTSRSS TTTEQLQFTI WDELIIFPIQ PLTCGTKLLY QHNLETLEND WGNLAKTYSL	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART HRAVDQVIK TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV LKGKLLDILH LHQWPALYPL	KLQKDRQVTD KLQKDRQVTD KLQLDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ KDSSLGLSKE IALELLDSKP	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSILLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV ALYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDR DKAFLWERRY ADQEVRSLAV	120 180 240 300 360 420 480 540 600 720 780 840 900
55 60	Protein Acc  1     MAQIFSNSGF TERKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KDPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNNHCLGSH CKEAMTRHPV KKLKRAVNLP NSGRSPTDCA FKPIQSKKVG QRKGPEALGK PSPAFDIIY TYCFKHPNCLP TWIEAISDDE	11    KECPFSHPEP   QDYDLMVFPE   LYFRPTIQRG   GQSPYFSYPL   SKVSNLQVSP   ERSTANCHLE   LQEVEVQNEE   EGFQLPVTFT   EHIQNCRKWD   EELLDSYHNQ   ESKKADVTSL   CSSKSVKEAW   TYKNFFYLIK   VSLPLCDFRR   TPQVDRSIIR   KILASAPNWK   LTDLLPQFVQ	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQFTI WDELIIFPIQ PLICIFKLLY QHNLETLENY ALKYEIYLNS	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH LKTKFPYTNH SAMCQNLART HRAVDQVIKA TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLLDILH LHQWPALYPL SLVQFLLSRA	KLQKDRQVTD KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ PGTVTKKGYV KDSSLGLSKE LALELLDSKP LGNIQIAHNL	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VIKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWEKRY ADQEVRSLAV YWLLKDALHD	120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020
<ul><li>55</li><li>60</li><li>65</li></ul>	Protein Acc  I	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD EELLDSYHNQ RSKTADVTSL QSSKSVKEAW TYKNFFYLIK VSLPLCDFRR TPQVDRSIIQ KILASAPNWK LTDLLPQFVQ LGALLSVGGK	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQPTI WDELIIFPIQ PLTCGTKLLY QHNLETLEND WGNLAKTYSL ALKYEIYLNS RLREELLKQT	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKEP ATVTRSQSLN LKTKFPYTHH IMQALCWVHD SAMCQNLART HRAVDQVIKA TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KKLQLAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ PGTVTKKGYV KDSSLGLSKE IALELLDSKF IALELLDSKF LGNIQIAHNL EKVRQASGSA	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKOPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWEKRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME	120 180 240 300 360 420 540 660 720 780 840 900 960 1020
<ul><li>55</li><li>60</li><li>65</li></ul>	Protein Acc  1	11    KECPFSHPEP   QDYDLMVFPE   LYFRPTIQRG   GQSPYFSYPL   SKVSNLQVSP    ERSTANCHLE   LQEVEVQNEE   EGFQLPVTFT   EHIQNCRKWD   EELLDSYHNQ   EELLDSYHNQ   EESLADVTSL   QSSKSVKEAW   TYKNFFYLIK   VSLPLCDFRR   TPQVDRSIIQ   KILASAPNWK   LTDLLPQFVV   LGALLSVGGK   CRLPLKPSLV	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQFTI WDELLIFPIQ PLICGTKLLY QHNLETLEND WGNLAKTYSL ALKYEIYLNG RLREELLKQT AKELNIKSCS	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA PFSSNAVPLK	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KLQLAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ FGTVTKKGYV KDSSLGLSKE IALELLDSKP LGNIQIAHNI EKVRQASGSA VTMVNADPLG	NORGFELSSS NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWEKRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG	120 180 240 300 420 480 540 600 660 720 780 840 900 960 1020 1080
<ul><li>55</li><li>60</li><li>65</li></ul>	Protein Acc  1	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD EELLDSYHNQ RSKTADVTSL QSSKSVKEAW TYKNFFYLIK VSLPLCDFRR TPQVDRSIIQ KILASAPNWK LTDLLPQFVQ LGALLSVGGK	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQFTI WDELLIFPIQ PLICGTKLLY QHNLETLEND WGNLAKTYSL ALKYEIYLNG RLREELLKQT AKELNIKSCS	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA PFSSNAVPLK	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KLQLAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ FGTVTKKGYV KDSSLGLSKE IALELLDSKP LGNIQIAHNI EKVRQASGSA VTMVNADPLG	NORGFELSSS NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWEKRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG	120 180 240 300 360 420 540 660 720 780 840 900 960 1020
<ul><li>55</li><li>60</li><li>65</li><li>70</li></ul>	Protein Acc  1	11    KECPFSHPEP   QDYDLMVFPE   LYFRPTIQRG   GQSPYFSYPL   SKVSNLQVSP   ERSTANCHLE   LQEVEVQNEE   EGFQLPVTFT   EHIQNCRKWD   EELLDSYHNQ   CSKSVKEAW   TYKNFFYLIK   VSLPLCDFR   TPQVDRSIIQ   KILASAPNWK   LTDLLPQFVQ   LGALLSVGGK   CRLPLKPSLV   QMIKIMDKIW	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIEND PGGEDTSRSS TTTEQLQFTI WDELIIFPIQ PLTCGTKLLY WGNLAKTYSL ALKYEIYLNS RLREELLKGI AKELNIKSCS LKEGLDLRMV	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA FFSSNAVPLK IFKCLSTGRD	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ FGTVTKKGYV KDSSLGLSKE IALELLDSKP LGNIQIAHNL EKVRQASGSA VTMVNADPLG RGMVELVPAS	NORGFELSSS NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSILLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV ALYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWERRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG DTLRKIQVEY	120 180 240 300 420 480 540 660 720 780 960 1020 1080 1140 1200
<ul><li>55</li><li>60</li><li>65</li><li>70</li></ul>	Protein Acc  1     MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KDPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNNHCLGSH CKEAMTRHPV KKLKRAVNLP NSGRSPTDCA FKPIQSKKVG QRKGPEALGK PSPAFDIIY TVCFKHPNCLP TWIEAISDDE VQFSTRYEHV RVQSFFQNK GVTGSFKDKP	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD EELLDSYHNQ RSKTADVTSL RSKTADVTSL QSSKSVKEAW TYKNFFYLIK VSLPLCDFRR TPQVDRSIIQ KILASAPNWK LTDLLPQFVQ LGALLSVGGK CRLPLKPSLV LAEWLRKYNP	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQFTI WDELIIFPIQ PLICTIKLLY QHNLETLEND ALKYEIYLNS RLREELLKGT AKELNIKSCS LKEGLDLRMV SEEEYEKASE	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART HRAVDQVIKA TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV LKGKLLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA FFSSNAVPLK IPKCLSTGRD	KLQKDRQVTD KLQKDRQVTD EKLLLDDSFE TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ PGTVTKKGYV KDSSLGLSKE LALELLDSKF LGNIQIAHNL EKVRQASGSA VTMVNADPLG RGMVELVPAS VATYVLGICD	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWEKRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG DTLRKIQVEY RHNDNIMLRS	120 180 240 300 420 480 540 600 720 780 840 900 960 1020 1080 11200 1260
<ul><li>55</li><li>60</li><li>65</li></ul>	Protein Acc  I      MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KDPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNMHCLGSH CKEAMTRHPV KKLKRAVNLP NSGRSPTDCA FKPIQSKVG ORKGPEALGK PSPAFDIIYT YCFKHPNCLP TWIEAISDDE VQFSTRYEHV RVQSFFQKNK EDLRQDMLAL GVTGSFKAKP TCHMFHIDFG	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD RSKTADVTSL QSSKSVKEAW TYKNFFYLIK VSLPLCDFRR TPQVDRSIIQ KILASAPNWK LTDLLPQFVQ LGALLSVGGK CRLPLKPSLV QMIKIMDKIN KFLGHAQMFG	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW KKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRQLLTF TEIRQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQFTI WDELIIFPIQ PHITCGTKLLY QHNLETLEND WGNLAKTYSL ALKYEIYLNS RLREELLKQT AKELNIKSCS LKEGLDLRMV SEEEYEKASE SFKRDRAPFV	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKEP KATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA FFSSNAVPLK IFKCLSTGRO NPIYSCAGCC LTSDMAYVIN	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KLQLAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ HGTVTKKGYV KDSSLGLSKE IALELLDSKP VROSGSA VTMVNADPLG RGMVELVPAS VATYVLGICD GGEKPTIRFQ	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWEKRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFRVG DTLRKIQVEY RHNDNIMLRS LFVDLCCQAY	120 180 240 300 360 420 660 720 780 840 900 900 1020 1080 1140 1260 1320
<ul><li>55</li><li>60</li><li>65</li><li>70</li></ul>	Protein Acc  1	11    KECPFSHPEP   CONTINUE     KECPFSHPEP   CONTINUE     KECPFSHPEP   CONTINUE	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW KKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQFTI WDELIIFPIQ PLTCGTKLLY QHNLETLEND WGNLAKTYSL ALKYEIYLNS ALKYEIYLNS RKPELIKQT AKELNIKSCS LKEGLDLRMV SEEEYEKASE SFKRDRAPFV GLPELTSIQD	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA PFSSNAVPLK IFKCLSTGRD NPIYSCAGCC LTSDMAYVIN LKYVRDALQP	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KLQLAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ PGTVTKKGYV KDSSLGLSKE IALELLDSKP LGNIQIAHNI EKVRQASGSA VTMVNADPLG RGMVELVPAS VATYVLGICR QGTKDFIFQ QTTDAEATIF	NORGFELSSS NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWERRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG DTLRKIQVEY RHNDNIMLRS LFVDLCCQAY FTRLIESSLG	120 180 240 300 360 420 660 720 780 960 1020 1080 1140 1260 1320 1380
<ul><li>55</li><li>60</li><li>65</li><li>70</li></ul>	Protein Acc  1	11    KECPFSHPEP QDYDLMVFPE LYFRPTIQRG GQSPYFSYPL SKVSNLQVSP ERSTANCHLE LQEVEVQNEE EGFQLPVTFT EHIQNCRKWD RSKTADVTSL QSSKSVKEAW TYKNFFYLIK VSLPLCDFRR TPQVDRSIIQ KILASAPNWK LTDLLPQFVQ LGALLSVGGK CRLPLKPSLV QMIKIMDKIN KFLGHAQMFG	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW KKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQFTI WDELIIFPIQ PLTCGTKLLY QHNLETLEND WGNLAKTYSL ALKYEIYLNS ALKYEIYLNS RKPELIKQT AKELNIKSCS LKEGLDLRMV SEEEYEKASE SFKRDRAPFV GLPELTSIQD	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA PFSSNAVPLK IFKCLSTGRD NPIYSCAGCC LTSDMAYVIN LKYVRDALQP	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KLQLAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ PGTVTKKGYV KDSSLGLSKE IALELLDSKP LGNIQIAHNI EKVRQASGSA VTMVNADPLG RGMVELVPAS VATYVLGICR QGTKDFIFQ QTTDAEATIF	NORGFELSSS NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWERRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG DTLRKIQVEY RHNDNIMLRS LFVDLCCQAY FTRLIESSLG	120 180 240 300 360 420 660 720 780 840 900 900 1020 1080 1140 1260 1320
<ul><li>55</li><li>60</li><li>65</li><li>70</li></ul>	Protein Acc  1	11    KECPFSHPEP   CONTINUE     KECPFSHPEP   CONTINUE     KECPFSHPEP   CONTINUE	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQFTI WDELIIFPIQ PLTCGTKLLY WGNLAKTYSL ALKYEIYLNS RLREELLKQI ALKYEIYLNS RLREELLKGCS LKEGLDLRMV SEEEYEKASE SFKRDRAPFV GLPELTSIQU LPSNDEPILS	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA FFSSNAVPLK IFKCLSTGRD NPIYSCAGCC LTSDMAYVI LKYVRDALQP PSPKTYSFRQ	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KLQLAFQKI TYSKQAAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ FGTVTKKGYV KDSSLGLSKE IALELLDSKP LGNIQIAHNL EKVRQASGSA VTMVNADPLG RGMVELVPAS VATYVLGICD GGEKPTIRFQ GGTDAEATIF DGRIKEVSVF	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV ALYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWERRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG DTLRKIQVEY RHNDNIMLRS LFVDLCCQAY FTRLIESSLG TYHKKYNPDK	120 180 240 300 360 420 660 720 780 960 1020 1080 1140 1260 1320 1380
<ul><li>55</li><li>60</li><li>65</li><li>70</li></ul>	Protein Acc  1     MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KDPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNNHCLGSH CKEAMTRHPV KKLKRAVNLP NSGRSPTDCA FKPIQSKKVG QRKGPEALGK PSPAFDIIY TYCFKHPNCLP TWIEAISDDE VQFSTRYEHV RVQSFFQNK TGHMFHIDFG NLIRKQTNLF SIATKFNFFI HYIYVVRILW	11    KECPFSHPEP   QDYDLMVFPE   LYFRPTIQRG   GQSPYFSYPL   SKVSNLQVSP   ERSTANCHLE   LQEVEVQNEE   EGFQLPVTFT   EHIQNCRKWD   ESLIDSYHNQ   ESLIDSYHNG   ESLIDSYHNG   ESLIDSYHNG   KILASAPNWK   LYKNFFYLIK   VSLPLCDFRR   TPQVDRSII    KILASAPNWK   LITDLLPQFVQ   LGALLSVGGK   CRLPLKPSLU   LAEWLRKYNP   KPLGHAQMFG   LNILLSLMIFG   LNILLSLMIFG   ENGLEPSFVF	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQFTI WDELIIFPIQ PLICTFKLLY QHNLETLEND ALKYEIYLNS RLREELLKQT AKELNIKSCS LKEGLDLRMV SEEEYEKASE SFKRDRAPFV GLPELTSIQU LPSNDEPILS RTFVEFQELH	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWHD SAMCQNLART HRAVDQVIKA TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA FFSSNAVPLK IFKCLSTGRD NPIYSCAGCC LTSDMAYVIN LKYVRDALQP PSPKTYSFRQ NKLSIIFPLW	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KKLQLAFQN TDMAKLFDKI VDNVEVLDHA RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ PGTVTKKGYV KDSSLGLSKE IALELLDSKE IALELLDSKE IALELLDSKE VTMVNADPLG RGMVELVPAS VATYVLGICD GGEKPTIRFQ QTTDAEATIF DGRIKEVSVF KLPGFPNRMV	NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKOPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWERRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG DTLRKIQVEY RHNDNIMLRS LFVDLCCQAY FTRLIESSLG TYHKKYNPDK LGRTHIKDVA	120 180 240 300 360 420 660 720 780 840 900 960 1020 1140 1290 1260 1320 1380 1380 1500
<ul><li>55</li><li>60</li><li>65</li><li>70</li><li>75</li></ul>	Protein Acc  1	11    KECPFSHPEP   QDYDLMVFPE   LYFRPTIQRG   GQSPYFSYPL   SKVSNLQVSP    ERSTANCHLE   LQEVEVQNEE   EGFQLPVTFT   EHIQNCRKWD   EELLDSYHNQ   EELLDSYHNQ   EELLDSYHNQ   EELLDSYHNQ   EELLDSYHNQ   TYKNFFYLIK   TPQVDRSIIQ   KILASAPNWK   LTDLLPQFVQ   LGALLSVGGK   CRLPLKPSLV   QMIKIMDKIW   KPLGHAQMFG   LNLLSLMIPS   LNLLSLMIPS	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV TEIRLQLLTF CDVSSTVEII TEIRLQLLTF VELALQIENQ PGGEDTSRSS TTTEQLQFTI WDELLIFPIQ WGNLAKTYSL ALKYEIYLNS RLREELLKOT AKELNIKSCS LKEGLDLRMV SEESYEKASE SFKRDRAPFV GLPELTSIQD LPSNDEPILS RTFVEFQELH VAECDLVCTF	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV IKGKLLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA PFSSNAVPLK IFKCLSTGRD KLYSTGRD KYVRDALQP PSPKTYSFRQ KKIVIFPLW FHPLLRDEKA	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KLQLAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVUD VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ PGTVTKKGYV KDSSLGLSKE IALELLDSKF IA	NORGFELSSS NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV AIYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWEKRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG DTLRKIQVEY FRHDDNIMLRS LFVDLCCQAY FTRLIESSLG TYHKKYWFDK LGSTHIRDVA SFSPTPGQIG	120 180 240 300 360 420 660 720 780 840 900 910 1020 1140 1260 1320 1380 1440 1560
<ul><li>55</li><li>60</li><li>65</li><li>70</li></ul>	Protein Acc  1    MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KOPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNNHCLGSH CKEAMTRHPV KKLKRAVNLP NSGRSPTDCA FKPIQSKKVG QRKGPEALGK FSPAFDIIYT YCFKHPNCLP TWIEAISDDE TGHMFHIDFG NLIRKQTNLF SIATKFNFFI KYIYVVRILW ARRKIELNSY GAVKLSISYR	11    KECPFSHPEP   QDYDLMVFPE   LYFRPTIQRG   GQSPYFSYPL   SKVSNLQVSPE   ERSTANCHLE   ERSTANCHLE   ERSTANCHLE   EGFQLPVTFT   EHIQNCRKWD   EELLDSYHNQ   EELLDSYHNQ   ESKSVKEAW   TYKNFFYLIK   VSLPLCDFRR   TPQVDRSIIQ   KILASAPNWK   LTDLLPQFVQ   KILASAPNWK   LTDLLPQFVQ   CGLPLKPSLV   QMIKIMDKIW   LAEWLRKYNPP   LAEWLRKYNPP   LNILSLMIPS   HNLAQLRFSG   EGQIEPSFVF   LQSLMNASTD   LQSLMNASTD	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRQLLTF VELALQIENF VELALQIENF VELALQIENF WDELIIFPIQ PLTCGTKLLY QHNLETLEND WGNLAKTYSL ALKYEIYLNS LKEELLKQT AKELNIKSCS LKEGLDLRMV SEEEYEKASE SFKRDRAPFV GLPELTSIQD LPSNDEPILS RTFVEFQELL RTFVEFQELL IKDLVTEDGA	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV LWTSSHTNSV LKYKFLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA PFSSNAVPLK IFKCLSTGRD NPIYSCAGCC LTSDMAYVIN LKYVRDALQP FSPKTYSFRQ NKLSIIFPLW NKLSIIFPLW DPNYVKTYL	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KLQLAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ PGTVTKKGYV KDSSLGLSKE IALELLDSKP LGNIQIAHNL EKVRQASGSA VTMVNADPLG RGMVELVPAS VATYVLGICD QTTDAEATIF QGTKEVSVF KLPGFPNRMV KLPGFPNRMV LPDNHKTSKR	NORGFELSSS NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV ALYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWERRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG DTLRKIQVEY RHNDNIMLES LFVDLCQAY FTRLIESSLG TYHKKYNPDK LGRTHIKDVA RSPSPTPGQIG KTKISRKTN	120 180 240 300 360 420 480 660 720 780 840 900 960 1020 1140 1200 1260 1320 1380 1440 1500 1620
<ul><li>55</li><li>60</li><li>65</li><li>70</li><li>75</li></ul>	Protein Acc  1    MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KOPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNNHCLGSH CKEAMTRHPV KKLKRAVNLP NSGRSPTDCA FKPIQSKKVG QRKGPEALGK FSPAFDIIYT YCFKHPNCLP TWIEAISDDE TGHMFHIDFG NLIRKQTNLF SIATKFNFFI KYIYVVRILW ARRKIELNSY GAVKLSISYR	11    KECPFSHPEP   QDYDLMVFPE   LYFRPTIQRG   GQSPYFSYPL   SKVSNLQVSP    ERSTANCHLE   LQEVEVQNEE   EGFQLPVTFT   EHIQNCRKWD   EELLDSYHNQ   EELLDSYHNQ   EELLDSYHNQ   EELLDSYHNQ   TYKNFFYLIK   TYPQVDRSIIQ   KILASAPNWK   LTDLLPQFVQ   LGALLSVGGK   CRLPLKPSLV   QMIKIMDKIW   KPLGHAQMFG   LNLLSLMIPS   LNLLSLMIPS	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRQLLTF VELALQIENF VELALQIENF VELALQIENF WDELIIFPIQ PLTCGTKLLY QHNLETLEND WGNLAKTYSL ALKYEIYLNS LKEELLKQT AKELNIKSCS LKEGLDLRMV SEEEYEKASE SFKRDRAPFV GLPELTSIQD LPSNDEPILS RTFVEFQELL RTFVEFQELL IKDLVTEDGA	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV LWTSSHTNSV LKYKFLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA PFSSNAVPLK IFKCLSTGRD NPIYSCAGCC LTSDMAYVIN LKYVRDALQP FSPKTYSFRQ NKLSIIFPLW NKLSIIFPLW DPNYVKTYL	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KLQLAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ PGTVTKKGYV KDSSLGLSKE IALELLDSKP LGNIQIAHNL EKVRQASGSA VTMVNADPLG RGMVELVPAS VATYVLGICD QTTDAEATIF QGTKEVSVF KLPGFPNRMV KLPGFPNRMV LPDNHKTSKR	NORGFELSSS NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV ALYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWERRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG DTLRKIQVEY RHNDNIMLES LFVDLCQAY FTRLIESSLG TYHKKYNPDK LGRTHIKDVA RSPSPTPGQIG KTKISRKTN	120 180 240 300 360 420 660 720 780 840 900 910 1020 1140 1260 1320 1380 1440 1560
<ul><li>55</li><li>60</li><li>65</li><li>70</li><li>75</li></ul>	Protein Acc  1    MAQIFSNSGF TRKKAQVYNK PILSPSFSAQ STEPIYLSLP KARTDLEITD KOPWDAVLLE TSSLPTGSSL NASVKVSIDI LQNNHCLGSH CKEAMTRHPV KKLKRAVNLP NSGRSPTDCA FKPIQSKKVG QRKGPEALGK FSPAFDIIYT YCFKHPNCLP TWIEAISDDE TGHMFHIDFG NLIRKQTNLF SIATKFNFFI KYIYVVRILW ARRKIELNSY GAVKLSISYR	11    KECPFSHPEP   QDYDLMVFPE   LYFRPTIQRG   GQSPYFSYPL   SKVSNLQVSPE   ERSTANCHLE   ERSTANCHLE   ERSTANCHLE   EGFQLPVTFT   EHIQNCRKWD   EELLDSYHNQ   EELLDSYHNQ   ESKSVKEAW   TYKNFFYLIK   VSLPLCDFRR   TPQVDRSIIQ   KILASAPNWK   LTDLLPQFVQ   KILASAPNWK   LTDLLPQFVQ   CGLPLKPSLV   QMIKIMDKIW   LAEWLRKYNPP   LAEWLRKYNPP   LNILSLMIPS   HNLAQLRFSG   EGQIEPSFVF   LQSLMNASTD   LQSLMNASTD	IP_002636.1  21    TRAKDVDKEE SDSQKRALDI QWPPGLPGPS TPATPFHPQG KSEDISKFDW RKVNGKSLSV MAAFCRSITK CDVSSTVEII TEIRQLLTF VELALQIENF VELALQIENF VELALQIENF WDELIIFPIQ PLTCGTKLLY QHNLETLEND WGNLAKTYSL ALKYEIYLNS LKEELLKQT AKELNIKSCS LKEGLDLRMV SEEEYEKASE SFKRDRAPFV GLPELTSIQD LPSNDEPILS RTFVEFQELL RTFVEFQELL IKDLVTEDGA	ALQMEAEALA DVEKLTQAEL TYALPSIYPS SLPIYRPVVS LDLDPLSKPK ATVTRSQSLN LKTKFPYTNH IMQALCWVHD SAMCQNLART TRGSLNPENP FAAHGISSNW ISQLPLESVL LWTSSHTNSV LWTSSHTNSV LKYKFLDILH LHQWPALYPL SLVQFLLSRA KLVQLLGGVA PFSSNAVPLK IFKCLSTGRD NPIYSCAGCC LTSDMAYVIN LKYVRDALQP FSPKTYSFRQ NKLSIIFPLW NKLSIIFPLW DPNYVKTYL	KLQKDRQVTD KLQKDRQVTD KLQKDRQVTD KLQLAFQN TDMAKLFDKI VDNVEVLDHE IRTTQLAKAQ RTNPGYLLSP DLNQVDVGSY AEDDETPVDL VRKICSALDG VQVSINQLTA VSNYEKYYLI HLTLFGILNQ PGTVTKKGYV KDSSLGLSKE IALELLDSKP LGNIQIAHNL EKVRQASGSA VTMVNADPLG RGMVELVPAS VATYVLGICD QTTDAEATIF QGTKEVSVF KLPGFPNRMV KLPGFPNRMV LPDNHKTSKR	NORGFELSSS NORGFELSSS TKKTPVLPVT GFNPRMPTFP ASTSEFLKNG EEKNVSSLLA GHISQKDPNG VTAQRNICGE VLKVCGQEEV NKHLYQIEKP VETLAITESV ALYDLLRLHA CSLSHNGKDL SSGSSPDSNK MERIVLQVDF DKAFLWERRY ADQEVRSLAV YWLLKDALHD RQVVLQRSME EEINVMFKVG DTLRKIQVEY RHNDNIMLES LFVDLCQAY FTRLIESSLG TYHKKYNPDK LGRTHIKDVA RSPSPTPGQIG KTKISRKTN	120 180 240 300 360 420 480 660 720 780 840 900 960 1020 1140 1200 1260 1320 1380 1440 1500 1620

Seq ID NO: 31 DNA sequence Nucleic Acid Accession #: CAT cluster

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					GAGGGCTGAG		840
25					AGGACAGTTT CGGAGGTGCC		900 960
23					TCAACAAGAT		1020
					TGCATCTTCA		1080
					ACCTGGAGAC		1140 1200
30					CCCGTCCTAG GAGCATTTGA		1260
	CTCTTAAGAA	CTATACATTT	GTATGATAAT	CCTCTGTCTT	TTGTGGGGAA	CTCAGCATCT	1320
					CAAGCATGGT TGACAGGTAC		1380 1440
					GGACTTTGGA		1500
35	AATAATATAA	GAGACCTTCC	AAGTTTTAAT	GGTTGCCATG	CTCTGGAAGA	AATTTCTTTA	1560
					AAGGCCTGAT		1620
					GTAGAGCTTT		1680 1740
					AGCTGAAAGA		1800
40					ATGCTTATCA		1860
					ATAACAGCCT TCACAAGCAC		1920 1980
					CAGGTGCTTT		2040
A.E.					GGTTCATTTT		2100
45					CTTGTACATC TCATGGGAAT		2160 2220
					CTGAATTTGG		2280
					TTTTCTCCTC		2340
50					CTGCAAAAGA CCCTTTCGGC		2400 2460
50					AATATTCTGC		2520
					TCACTGTAAC		2580
					CTAAGCTATA TTAAGCATGT		2640 2700
55					TTTCATTTGC		2760
					CTCTGATATT		2820
					CAAAGTTTAA TTTCAGTTTC		2880 2940
					GCATGTACTC		3000
60						AGTATCATGC	
						CCAAAGACCT AGATGAAGAA	
						CTGCTTCTAC	
<b>C E</b>						TAAAGACTGA	
65						GAGTGAACCC ACTTAGAAGA	
						GCCTAAATTA	
	TAAATTGGTG	AAAAATGCAA	TGTCCAAGCA	ATGTATGATC	TGTTTGAAAC	AAATATATGA	3540
70						GATCCATAAG TTAATATTTT	
70						GCTAATTTTA	
	CCTAATGTTT	CATCCTTAAT	CTCAGGACAA	CTTACTGCAG	GGCCAAAAA	GGGACTGTCC	3780
`						TTGCCATCCT CTGAAGATGT	
75						GTTTTCAGTC	
. –	ATTATACATT	GCTTTGGTCC	AATCAGTAAT	TTTTTCTTAA	GTGTTTTGTG	ATTACACTAC	4020
						AAACTACTAA	
						GTTCTCATTA	
80	TAGCAATAGC	TTGGATTATA	TAGAAAGTAA	ACTGTGGTCA	ATACTTGCAT	TTAATTAGAC	4260
						GCTGGATTAT	
						TTTTTTATAA ATGTTATTAA	

	TAAAAATAGA	AGAAGAAAGA	ATAAAGCTTA	GTCCTGTGTC	TTTAAAAATT	ATTTTAAAAA	4500
	CTTGATTCCC	ATCTATGGGC	TTTAGACCTA	TTACTGGGTG	GAGTCTTAAA	GTTATAATTG	4560
	TTCAATATGT	TTTTTGAACA	GTGTGCTAAA	TCAATAGCAA	ACCCACTGCC	ATATTAGTTA	4620
_	TTCTGAATAT	ACTAAAAAAA	TCCAGCTAGA	TTGCAGTTTA	ATAATTAAAC	TGTACATACT	4680
5				ATTATTTTTA			4740
•				ACCTCCTAAA			4800
				TGCATTGTAA			4860
				TCAAAATCAT			4920
				TGTAAGGCCA			4980
10							
10				AAATATTAAA			5040
				GTTATTGAAA			5100
				TTAATCTTTG			5160
	AATATTGTAC	TCAGTGTTTT	GAATTATTAA	AGTTTCTAGA	AAGCAAAAAA	A	
1.							
15	Seq ID NO:	36 Protein	sequence				
	Protein Acc	cession #: N	IP 060960.1				
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20	MDGDLGLLCE	LALCILICEAC	DCCAADDI.CA	APCSCDGDRR	VDCSGKGLTA	UDECL SARTO	60
20				NDLSFIHPKA			120
				GLVQLRHLWL			180
				NKIRGLSQHC			240
25				NPLLRTIHLY			300
25				ISSIPNNLCQ			360
				LRILDLSRNL			420
	SFNELTSFPT	EGPNGLNQLK	LVGNFKLKEA	LAAKDFVNLR	SLSVPYAYQC	CAFWGCDSYA	480
	NLNTEDNSLQ	DHSVAQEKGT	ADAANVTSTL	ENEEHSQIII	HCTPSTGAFK	PCEYLLGSWM	540
				PSSKLFIGLI			600
30	SWGRFAEFGI	WWETGSGCKV	AGFLAVESSE	SAIFLLMLAT	VERSLSAKDI	MKNGKSNHLK	660
				PLCLPFPTGE			720
				WLIFTNCIFF			780
				DWKLLKRRVT			840
				SCKHLIKSHS			900
35				FYQSRGFPLV			300
55	TQSARSDIAD	EEDSE A2D22	DQVQACGRAC	FIQSKGFPDV	RIMINDPRVK	D	
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	1	1	1	1	1	]	
	 ATGCTGCGAG	 CCGCAGTGAT	1	 ATCAGGACCT	 GGCTCGCGGA	]	60
4.5			 CCTGCTGCTC	1		] GGGCAACTAC	60 120
45	CCCAGTCCCA	TCCCGAAATT	CCTGCTGCTC CCACTTCGAG	 ATCAGGACCT	CTGTGCCCGA	 GGGCAACTAC AGTCGTCCTG	
45	CCCAGTCCCA AACCTCTTCA	TCCCGAAATT ACTGCAAAAA	 CCTGCTGCTC CCACTTCGAG TTGTGCAAAT	ATCAGGACCT TTCTCCTCTG	CTGTGCCCGA TTCAAAAGAT	GGGCAACTAC AGTCGTCCTG TTTGGACAGG	120
45	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA	TCCCGAAATT ACTGCAAAAA GATACGATGT	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG	120 180
45	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA	 ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG	120 180 240
	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA GACTTGGAAA	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC	120 180 240 300
	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTTGA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA GACTTGGAAA TCGGATGCAT	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT	120 180 240 300 360 420
45 50	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTTGA TTGAACAGCA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA GACTTGGAAG TCGGATGCAT CGTGCATGAT	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT	120 180 240 300 360 420 480
	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTTGA TTGAACAGCA CACCCAGATG	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG	CCTGCTGCTC CCACTTCGAC TTGTGCAAAT CCGCCTGAGA GACTTGGAAA TCGGATGCAT CGTGCATGAT GTACGGCATC	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCA	CTGTGCCGA TTCAAAGAT GAGGTGCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG	120 180 240 300 360 420 480 540
	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTTGA TTGAACAGCA CACCCAGATG GATCTGCATA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA GACTTGGAAA TCGGATGCAT CGTGCATGAT GTACGGCATCA GACAAGCAG	I ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCA GCCTGCAACC	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA	GGGCAACTAC AGTCGTCCTG TTTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGGT	120 180 240 300 360 420 480 540
	CCCAGTCCCA AACCTCTTCAA GTGCTGTCAA AGAATATCTA ATCAGGATGT CTGAACTGA TTGAACAGCA CACCCAGATG GATCTGCATA TACACGGTTG	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AAGACATCAT	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACCATTGAAA TCGGATGCAT CGTGCATGAT GTACGGCATC GGACAAGCAG ATTATTCTGG	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAC GATTCACGCT GAGAAGTTGT GTGACTCACCG GCCTGCAACC GATGACAATG	CTGTGCCCGA TTCAAAAGAT GAGGTGCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGAT CACATGACT	120 180 240 300 360 420 480 540 600 660
50	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCAGGATGT CTGAACTTGA TTGAACAGCA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AAGACATCAT ATATCCCTCA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAAA TCGGATGCAT CGTGCATGAT GTACGGCATC GGACAAGCAG ATTATTCTGG GTTCACTTTC	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CCACTCACCA GCCTGCAACC GCTTGCAACCA CATGGACAATG CTGGGAAGGA	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CGATTACTAG	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGAT CCACATGACT CAAGGAGGTG	120 180 240 300 360 420 480 540 600 660 720
	CCCAGTCCCA AACCTCTTCA AGAATATCTA ATCACGATGT CTGAACTGGA CTGAACTGGA CACCCAGATG GATCTGCATA TACACGGTTG GAGGGCTGC TATTTCTACA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AGGACATCAT ATATCCCTCA 'CAGGTTCCTA' 'CAGGTTCCTA'	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA GACTTGGAAA TCGGATGCAT GTGCATGAT GTACGGCATC GGACAAGCAG ATTATTCTGG GTTCACTTTC CATACGCCTG	I ATCAGGACCT TTCTCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GTGACTGTGG CGACTCACCA GCCTGCAACC GATGACAATG CTGGGAAGGA ATACTGAAGT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CGATTACTAG TCCAGGTTCA	GGGCAACTAC GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGGT CCACATGACT CAAGGAGGTG GAGGGAAGTT	120 180 240 300 360 420 480 540 600 720 780
50	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTGCA CACCCAGATG GATCTGCATA TACACGGTTG GAGGGCTGC GAGGGCTGC TATTTCTACA AACAGCTACC	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AAGACATCAT ATATCCCTCA TCAGGTTCCTA TTGTGCAAGT	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACTTGGAAA TCGGATGCAT GTACGGCATC GTACAGCAT GTACTGCATC GTACTGCATC GTACTCCCTC CATACCCCTC CTACTGGCCT	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CCGACTCACCA GCCTGCAACC GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CGATTACTAG TCCAGGTTCA CCACTATTAC	GGGCAACTAC AGTCGTCCTG AGTCGTCCTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGGT CCACATCACT CAAGGAGGTG GAGGAGGTT CAGGGAGGTT CTCTTGGATA	120 180 240 300 360 420 480 540 600 660 720 780 840
50	CCCAGTCCCA AACCTCTTCA AGCATGTCAA AGAATATCTA ATCAGGATGT CTGAACTTGA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACAGCTACC CCCTTTTGGA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AAGACATCAT ATATCCCTCA TTGTGCAAGT TTGTACAGTT TGAACTATGA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA TCGGATGCAT GTGCATGAT GTACGGCAT GTACAGCAG ATTATTCTGG GTTCACTTTC CATACGCCTG CTACTGGCAT TTCCTCTGCA	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCA GCCTGCAACC GATGACAACC CATGGAAGGA ATACTGAAGT ACTGGAAGGT ACCGGAAGGT ACCGGGAGGTGA	CTGTGCCCGA TTCAAAAGAT GAGGTGCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CGATTACTAG TCCAGGTTCG CCACTATTAC CAATTGGCTT	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGGT CCACATGACT CAAGGAGGTG GAGGGAAGTT CTCTTGGATA AACTTCAATG	120 180 240 300 360 420 480 540 660 720 780 840 900
50	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTTGA CACCCAGATG GATCTGCATA TACACGGTTG GAGGGGCTGC TATTTCTACA AACAGCTACC TCGTTTTGGA CTCATCTGA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTT GAACGGTGCG AATTCCCTAT AAGACATCAT ATATCCCTCA 'CAGGTTCCTA TTGTGCAAGT TGAACTATGA CCACCATCGA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA GACTTGGAA TCGGATGCAT GTACAGCAT GTACAGCAT GTACAGCAT GTACAGCAT GTACAGCAT GTACAGCAT GTACAGCTT CATACGCCT CTACTGCA CTCACATCTG CTCACATCTG	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCA GCCTGCAACC GCTGCAACC GTGGACGATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGG	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CGATTACTAG TCCAGGTTCA CCACTATTAC CAATTGCCTT TCCCCAACAT	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATCGT CCACATGACT CAAGGAGGTG GAGGGAAGTT CTCTTGGATA AACTTCAATG	120 180 240 300 360 420 480 540 600 660 720 780 840 900 960
50 55	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTAGCA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACAGCTACC TCGTTTTGGA ACCCTCTGA AAGGCCATTG	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AGGACTATCATA ATATCCCTCA TCAGGTTCCTA TCAGGTTCCTA TCAGGTTCCTA TCAGGTTCCTA AGACTATGA CCACCATCGA ATATCTATAT	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA GACTTGGAAA TCGGATGCAT GTACAGCAT GTACAGCAT GTACAGCAT GTACAGCTT CCATACGCCT CCATACGCCT TTCCTCTGCA CTCACATCTG CCTCGTGTGC CCTCGTGTGC	I ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACCT GTGACTGTGG GCACTCACCA GCCTGCAACC GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT TCCAGGTTCA CCACTATTAC CCACTATTAC TCCCCAACAT TGCTCCTGTC	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGGT CCACATGACT CAAGGAGGTG GAGGGAAGTT CTCTTGGATA AACTTCAATG TTCCTGGATA CTCTTGGATA CTCTTGGATA CTCTTGGATA CTCTTGGATA CTCTTGGATA CTCTTGGATA CTCTTGGATA CTCTTGGATA CTTCTTGGATA CTTCTTGGATA CTTCTTGGATA CTTCTTGGATA CTTCTTGGATA CTTCTTGGATA CTTCTTGGATA CTTGCTGGAG	120 180 240 300 360 420 480 540 660 720 780 840 900 960 1020
50	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTGCA TTGAACAGCA CACCCAGATG GATCTGCATA TACACGGTTG GAGGGCTGC TATTTCTACA AACAGCTACC TCGTTTTGGA CTCATCCTGGA AAGGCCATTG TATGTCTACA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT AAGACATCAT ATATCCCTAT ATATCCCTAT ATATCCCTAT TTGTGCAAGT TGAACTATGA CCACCATCGA ATATCTATAT TCAACTATCT TCAACTATCT	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA TCGGATGCAT CGTGCATGAT GTACGGCATC GGACAAGCAG ATTATTCTGG GTTCACTTTC CATACGCCTT CTACTGGCCT TTCCTCTGCA TTCCTCTGCA TTTCTGGTGTGC TTTCTTACAGT TTTCTACAGT	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGAATCTCAG GATCACAGCT GAGAAGTTGT GTGACTGTGG CGACTCACCC GATGACAACC GATGACAACC GATGACAACC GATGACAACC GATGACAACC GATGACAACC GATGACAACT CCGGGATAAGC CCGGGATAAGC CTGAGGACCT CCAGGGACCT CCAGGACCT CCAGGGACCT CCAGGGACCT CCAGGGACCT CCAGGACCT CCAGGACCT CCAGGACCT CCAGGACCT CCAGGACCT CCAGGACCT CCAGGACCT CCAGGACCT CCAGGACCT CCAGGACCC CCAGGGACCC CCAGGGACC CCAGGGACCC CCAGGGACCC CCAGGGACCC CCAGGGACC CCAGGGACCC CCAGGGACC CCAGGGACCC CCAGGCACC CCAGCACC CCAGGCACC CCAGCC CCAGCC CCAGCC CCAGCC CCAGCC CCAGCC CCAGCC CCAGCC CCACC	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CGATTACTAG TCCAGGTTCA CCACTATTAC CAATTGGCTT TCCTCCCAACAT TGGTCCTGTC GGCGCCAGCC	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGTGCCTGT GTTTCAGCTT TTTTCACGTT TTGTTCCCTG GAGCTATGAT CCACATGACT CAAGGAGGTG GAGGGAAGTT CTCTTGGATA AACTTCAATG TTCCTGTATC CTTGGAGT TAGGCGACAC	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020
50 55	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTTGA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACAGCTACC TCGTTTTGGA CTCATCCTGA AAGGCATTC TATGTTTACA AAGAGCCATTG	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AAGACATCAT ATATCCCTA 'CAGGTTCCTA TTGTGCAAGT TGAACTATGA CCACCATCGA ATATCTATAT TCAACTATAT TCAACTATCT GAAGAGTCAT	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA TCGGATGCAT GTGCATGAT GTACGCAT GTACAGCAT GTACAGCAT GTACAGCAT CCTACAGCCTG CTACTGCA CTACAGCCTG CTACTGCA CTCCTGCA CTCACATCTG CTTCCTGCA CTCACATCTG CTTCCTGCAGT TTCCTCTGCAT TTCCTACAGCT TTCCTACAGT TGCCCGCTAC	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCA GCCTGCAACC GATGACAACC CTGGAAGGA ATACTGAAGG ATACTGAAGT TCGTCATACCTCA GCCAGGGTGA CCGGATAAGC CCGGATACCC CCAGGACCTC CCGCTACCAGC	CTGTGCCCGA TTCAAAAGAT GAGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC CTGATGGTAGA GGAACGCCAT CCAGTATACTAG TCCAGGTTCA CCACTATTAC CAATTGCCTT TCCCCCAACAT TGTTCCTGTC GGCGCCAGCCC AAGTGGTGGT	GGGCAACTAC AGGCCACC TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGGT CCACATGACT CAAGGAGGTG GAGGGAAGTT CTCTTGGATA AACTTCAATG TTCCTGTATC CTTGCTGGAG TAGGCGACA AGGAAACGTG	120 180 240 300 360 420 480 540 600 660 720 780 840 900 960 1020 1080 1140
50 55	CCCAGTCCCA AACCTCTTCA AGAATATCTA ATCACGATGT CTGAACTTGA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACCTACC TCGTTTTGGA CTCATCTGA AAGGCCATTG TATTCTACA AAGGCCATTG CTCATCCTGA AAGGCCATTG TATTCTACA CCCAGATGCC CAGGATGGC	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTT GAACGGTGCG AATTCCCTAT ATATCCCTCA 'CAGGTTCCTA TTGTGCAAGT TCGAACTATGA CCACCATCGA ATATCTATAT TCAACTATCT CGAAGGTCAT TCAACTATCT TCAACTATCT TGAACTATCT TCAACTATCT TGAACGATCAT TCAACTATCT TGAACGATCAT TCAACTATCT TGAAGAGTCAT TGAATTAACGT	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA TCGGATGCAT GTACGCATG GTACAGCAT GTACGCATC GTACATCTC GATACGCCTG CTACTCTCC CTACTCTCC CTACTCTCC CTCCTGCA CTCACATCTG CTCCTGTGCA CTCCTCTGCA CTCCTCTGCA CTCCTCTGCA CTCCCGCTAC CTCCCGCTAC GGAAGACGGA	I ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCA GCCTGCAACC GATGACAACG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGGATAAGC TTGTTCTTTC CGAGGACCTC CGCTACCAGC GTCAGCTCTC	CTGTGCCCGA TTCAAAAGAT TGAGGTGCCCC AAATGAATAT TAGCATACTA AGGATCCCTA AGGATCCCTA CGATCAGCAGC TGTGGTAGA GGAACGCCAT CCACTATACTAG CCACTATTAC CCACTATTAC CAATTAGCTT TCCCCAACAT TGTTCCTGTC GGCGCCAGCG TCCCCATCAC TCCCCATCAC	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACCTT GTTTCAGCTT TTGTTCCCTG GAGCTACTT TGTTTCAGCTT CCACATGACT CCACATGACT CCACATGACT CTCTTGGATG TCTTTGATT CTCTTGGATG TCTCTTGATT CTCTTGATG TCCTGTATC CTTGCTGAGG TAGGCGACAC AGGAAACCTG CCCAGCGCAG	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020
50 55	CCCAGTCCCA AACCTCTTCA AGAATATCTA ATCACGATGT CTGAACTTGA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACCTACC TCGTTTTGGA CTCATCTGA AAGGCCATTG TATTCTACA AAGGCCATTG CTCATCCTGA AAGGCCATTG TATTCTACA CCCAGATGCC CAGGATGGC	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTT GAACGGTGCG AATTCCCTAT ATATCCCTCA 'CAGGTTCCTA TTGTGCAAGT TCGAACTATGA CCACCATCGA ATATCTATAT TCAACTATCT CGAAGGTCAT TCAACTATCT TCAACTATCT TGAACTATCT TCAACTATCT TGAACGATCAT TCAACTATCT TGAACGATCAT TCAACTATCT TGAAGAGTCAT TGAATTAACGT	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA TCGGATGCAT GTACGCATG GTACAGCAT GTACGCATC GTACATCTC GATACGCCTG CTACTCTCC CTACTCTCC CTACTCTCC CTCCTGCA CTCACATCTG CTCCTGTGCA CTCCTCTGCA CTCCTCTGCA CTCCTCTGCA CTCCCGCTAC CTCCCGCTAC GGAAGACGGA	I ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCA GCCTGCAACC GATGACAACG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGGATAAGC TTGTTCTTTC CGAGGACCTC CGCTACCAGC GTCAGCTCTC	CTGTGCCCGA TTCAAAAGAT TGAGGTGCCCC AAATGAATAT TAGCATACTA AGGATCCCTA AGGATCCCTA CGATCAGCAGC TGTGGTAGA GGAACGCCAT CCACTATACTAG CCACTATTAC CCACTATTAC CAATTAGCTT TCCCCAACAT TGTTCCTGTC GGCGCCAGCG TCCCCATCAC TCCCCATCAC	GGGCAACTAC AGGCCACC TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGGT CCACATGACT CAAGGAGGTG GAGGGAAGTT CTCTTGGATA AACTTCAATG TTCCTGTATC CTTGCTGGAG TAGGCGACA AGGAAACGTG	120 180 240 300 360 420 480 540 600 660 720 780 840 900 960 1020 1080 1140
<ul><li>50</li><li>55</li><li>60</li></ul>	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTGCA TTGAACAGCA CACCCAGATG GATCTGCATA TACACGGTTG GAGGGCTGC TATTTCTACA AACAGCTACC TCGTTTTGGA CTCATCCTGGA TATGTCTACA AAGGCCATTG TATGTCTACA AGGAGACCCC CAGGATGGCC CCCCCTGG CTGGCCACCT	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT AAGACATCAT ATATCCCTAT ATATCCCTAT TGTGCAAGT TGAACTATGA CCACATCGA ATATCTATAT TCAACTATCT TCAACTATCT GAAGAGTCAT TCAACTATCT GAAGAGTCAT TCAACTATCT CAAGCACGGA COGAAAGCCT CAGAAAGCCCGCA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA TCGGATGCAT CGTGCATGAT GTACGGCAT GTACGGCAT GTACGGCAT CCTACATCTG CTACATCTG CTCACATCTG CCTCACATCTG CCTCTGCAG TTCCTCTACAG TTTCTACAGT TGCCCGCTAC GGAAGACGGA AAGCCTCGG CAGCCCACTC	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAC GATCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCC GATGACAACC GATGACAACC GATGACAACC GATGACAACC GATGACAACC GATGACAACT CCGGATAAGC TCGGATAAGC TCGTTCCTCA CCAGGACTCCC CCTACACCC TCTTTGACGT TCTTTTGACGT ACTTCTCTCT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCACTATTAC CCACTATTAC CCACTATTAC CGCCCAACAT TGGTCCTGTC GGCGCCAGCC AAGTGGTGT TCCCCATCAC CCACCTCCGA CCACCTCCGA CCACCTCCGA CCACCTCCGA CCACCTCCGA CAGCCCAGGC CAGCCCAGGC CAGCCCAGGC	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGTGCCTGT GTGTCCTGT GTTTCAGCTT TTGTTCCCTG GAGCTACTT CACATGACT CACATGACT CACATGACT CACATGACT CATGACT CACATGACT CATGACT CACATGACT CACATGACT CACATGACT CACATGACT CACATGACT CACATGACT CCTGCAGCCAG CCCCCTGGCCAG CCCCCTGGCCAG CCCCCTGGCCCAG	120 180 240 300 360 420 540 600 660 720 780 840 900 960 1020 1080 1140 1200
<ul><li>50</li><li>55</li><li>60</li></ul>	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTGCA TTGAACAGCA CACCCAGATG GATCTGCATA TACACGGTTG GAGGGCTGC TATTTCTACA AACAGCTACC TCGTTTTGGA CTCATCCTGGA TATGTCTACA AAGGCCATTG TATGTCTACA AGGAGACCCC CAGGATGGCC CCCCCTGG CTGGCCACCT	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT AAGACATCAT ATATCCCTAT ATATCCCTAT TGTGCAAGT TGAACTATGA CCACATCGA ATATCTATAT TCAACTATCT TCAACTATCT GAAGAGTCAT TCAACTATCT GAAGAGTCAT TCAACTATCT CAAGCACGGA COGAAAGCCT CAGAAAGCCCGCA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA TCGGATGCAT CGTGCATGAT GTACGGCAT GTACGGCAT GTACGGCAT CCTACATCTG CTACATCTG CTCACATCTG CCTCACATCTG CCTCTGCAG TTCCTCTACAG TTTCTACAGT TGCCCGCTAC GGAAGACGGA AAGCCTCGG CAGCCCACTC	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACCA GCATGCACCA GCCTGCAACC GATGACAATG CTGGGAAGTGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG CCAGGATAGC TTGTTCTTTG CCAGGATCACC GCTACCAGC GTCAGCTTC TCTTTGACGT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCACTATTAC CCACTATTAC CCACTATTAC CGCCCAACAT TGGTCCTGTC GGCGCCAGCC AAGTGGTGT TCCCCATCAC CCACCTCCGA CCACCTCCGA CCACCTCCGA CCACCTCCGA CCACCTCCGA CAGCCCAGGC CAGCCCAGGC CAGCCCAGGC	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGTGCCTGT GTGTCCTGT GTTTCAGCTT TTGTTCCCTG GAGCTACTT CACATGACT CACATGACT CACATGACT CACATGACT CATGACT CACATGACT CATGACT CACATGACT CACATGACT CACATGACT CACATGACT CACATGACT CACATGACT CCTGCAGCCAG CCCCCTGGCCAG CCCCCTGGCCAG CCCCCTGGCCCAG	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020 1080 1140 1200 1260 1320
50 55	CCCAGTCCCA AACCTCTTCA AGAATATCTA ATCAGATGT CTGAACTTGG CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACCTACC TCGTTTTGGA CTCATCCTGA AAGGCCATTG TATTCTACA AAGGCCATTG CATCCCTGG CAGGATGGCC CAGGATGGCC CAGGATGGCC CAGGATGGCC CTGGCCACCT ACTGCAGAAA	TCCCGAAATT ACTGCAAAA GATACGATGA TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AGACATCAT ATATCCCTCA 'CAGGTTCCTA TTGTGCAAGT TCAACTATGA ACACTATGA ACACTATGA ATATCTATAT TCAACTATCT GAAGAGTCAT TGAACTATCT TGAACTATCT GAAGAGCCGGA CCGAAAGCC GCCGAAGCCGGA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA TCGGATGCAT GTACGGCAT GTACGGCAT GTACGGCAT GTACAGCCTG GTACAGCCTG CTACATCTC CATACGCCTG CTACATCTGC CTCCTGTGA TTCCTCTGCA TTCCTCTGCA TTCCTGCAGAGCCTC TTCTACAGCT TTCCTGCTACACT TTCCTCGCTAC CCTCGTGTGC AGACCTCC CGAAGACCGCAC CGAAGACCGCT CCACCTCC CCTCCCCTCC	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCA GCCTGCAACC GATGACAACC GATGACAACC GATGACAACC GATGACAACT ACTGTCATTGAG CCAGGGTGA CCGGATAAGC TTGTTCTTTG CCAGGACCTC CCCTACCAGC GTCAGCTCT ACTTCTCTCTC ACCTCCAGAGC ACCTCAGAGC ACCTCAGAGC ACCTCAGAGC	CTGTGCCCGA TTCAAAAGAT TGAGGTCCCCGA AAATGAATAT TAGCATACTA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT TCCAGGTTCA CCACTATTAC CCACTATTAC TCCCCAACAT TCTTCCTGTC GGCGCCAGCC AAGTGGTGGT TCCCCATCAC CCACTCCGA CAGGCCAGCC AAGTGGTGGT TCCCCATCAC CCACCTCCGA CAGGCCAGCC AGGCCAGGC AGGCCAGGC AGGCCAGGC	GGGCAACTAC GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGGT CCACATGACT CCACATGACT CTCTTGGATA CTCTTTGGATA TCCTTGCTGAG TCCTGCTGAG TCCTGCTGAG CCAGGCCAG CCAGGCCAG CCCCCTGGC CAGCCTATGCT	120 180 240 300 360 420 600 660 720 780 840 900 1020 1080 1140 1200 1260
<ul><li>50</li><li>55</li><li>60</li></ul>	CCCAGTCCCA AACCTCTTCA AGAATATCTA ATCACGATGT CTGAACTGGA CACCCAGATG GATCTGCATA TACACGGTTG TACACGGTTG TATTTCTACA AACAGCTACC TCGTTTTGGA CTCATTCTACA AAGGCCATTG TATTTCTACA AAGGCCATTG CTCATTCTACA AGGAGCCCC CAGGATGGCC GCCCCCTGG CTGGCCACCT ACTGGAAAA GTTCGCTTTA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGACGTGCG AATTCCCTAT AGACATCAT ATATCCCTCA 'CAGGTTCCTA TTGTGCAAGT TGAACTATGA CCACCATCGA ATATCTATAT TCAACTATCT GAAGAGTCAT GAAGAGTCAT GAAGAGTCAT GAAGAGTCAT GAAGAGTCAT GAAGAGTCAT GAAGAGTCAT CAACTATCT GAAGAGTCAT CAACTATCT GAAGAGTCAT CAACTATCT GAAGAGTCAT CAACTATCT GAAGAGTCAT CAAGCCCGGA ATGCTTACCAA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA TCGGATGCAT GTACGGCAT GTACGGCAT GTACGGCAT CCTCACTTC CATACGCCT CTACACTCT CCTCTGCA CTCACATCT CTCCTGCA CTCACATCT CCTCGCAT CTCACATCT CCTCGCAT CTCACATCT CCTCGCAT CTCACATCT CCTCGCAT CCTCGCAT CCTCACATCT CCTCGCAT CCTCACATCT CCTCGCAT CCTCACATCT CCTCCCTCC CGCACACC CAGCCCACT CGCACACC CAGCCCACT CTCTCCCCTCC CGCTGATGAC	I ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAGG GATTCACGCT GAGAAGTTGTG GTGACTGTGG CGACTCACCA CCTGCAACC GATGACAACC GATGACAACC CTGGGAAGGA ATACTGAAGT ACTGTCCTCA CCGGGATAAGC CTGGATACAGC CTGCAACC GTCAGCTCTC TCTTTGACGT ACTTCTCTCT TCTTTTGACGT ACTTCTCTCT ACTTCTCTCAGAGC AGTATTTTC	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA GGATCCCTGA CTACAGCAGC TGGTGGTAGA TCATACTAG TCAGGTTCA CCACTATTAC CAATTACTAG TCCCAACAT TGTTCCTGTC GGCGCCAGCC AAGTGGTGGT TCCCCATCAGC CAACTCCGA CCACCTCCGA CCACCTCCGA CCACCTCCGA CCAGCCCAGC	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGT CCACATGACT CCACATGACT CTCTTGGATG TCTCTGGATG TCTCTGGATG TCTCTGGATG TCTCTGATG TCCTGTATC CTTGCTGGAG TAGGCGACAC CCCAGCCAG CCCCTGGCC CCACCTGGCC CCACCTGGCC CCACCTGTT CCAGCACCGT CCGCAACCGT CCGCAACCGT	120 180 240 300 360 420 540 600 660 720 780 840 900 960 1020 1140 1200 1260 1380 1440
<ul><li>50</li><li>55</li><li>60</li></ul>	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTAGCA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACAGCTACC CGGTTTTGGA CTCATCCTGA AAGGCCATTG TATGTCTACA AGGAGCCCCCCGG GCCCCCTGG CTGGCCACCT ACTGGAAGACCC CTGGCCACCT ACTGGAAGACCC GTGGCCACCT ACTGGAAGACCC GTCGGCACCT ACTGCAAGCCC	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AGGACTATCAT ATATCCCTAT TCAGCTTCCTA TTGTGCAAGT TCAACTATCA CCACCATCGA ATATCTATAT TCAACTATCT GAAGAGTCAT TCAACTATCT GAAGAGTCAT CCACCACGA ATATCTATAT TCAACTATCT GAAGAGTCAT CCACCACGA ACCCCGGA CCGAAAGCCT GCCTGAGCGA ATGGTTCCA ATGGCCATGG	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA TCGGATGCAT GTGCATGAT GTGCATGAT GTACGGCAT GTACGGCAT GTACTGCA ATTATTCTGG GTTCACTTTC CATACGCCT CTACTGGCC TTCCTCTGCA TTCCTCTGCA TTCCTGTGTGC TTCCTGTGTGC TTCCTGTGTGC TTCCCCTCC GGAAGACGGA AAGCCTCGGTT CAGCCCTC CTCCCCTCC CGCTGATGAC GGTTAATCCCT TCTCCCCTCC CGCTGATGAC TCTCCCCTCC CGCTGATGAC TGTTACCCAT	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATCTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCA GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG CGAGGACCTC GTCAGCTCTC TCTTTGACTGT ACTTCTCTCT ACCTCAGAGG ACTTCTCTCT ACCTCAGAGG ACTTCTCTCT GACCATGAAGA GACATTTTC GACCATGAAGA GACATGAAGA GACATGAAGA GACATGAAGA GACATGAAGA GACATGAAGA GACATGAAGA GACATGAAGAC GACAATGAAGAC GACAATGAAAC GACAATGAAC GACAATGAAAC GACAATGAAC GACAATGAAAC GAC	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT TCCAGTTCA CCACTATTAC CCACTATTAC CCACTATTAC GGCGCCAGCC AAGTGGTGGT TCCCCGACCT TCCCCACCT CCACCTCCGA AGGCCCGGC AGGCCAGCC CAGCCCCGAA CTACCGAAT ATTCCAATGA	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACT GTGCACACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGACT CAAGGAGGTG CACATGACT CATTACATG CTCTTGGATA AACTTCAATG CTCTGGATA TTCCTGTATC CTTGCTGGAG AGGAACCTG CCCAGCCCAG	120 180 240 300 360 420 600 660 720 780 840 900 1020 1140 1260 1320 1320 1440 1500
<ul><li>50</li><li>55</li><li>60</li></ul>	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTGCA TTGAACAGCA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACAGCTACC TCGTTTTGGA AAGGCCATTG TATGTCTACA AGGAGCCCC CAGGATGGC CTCGCCCTGG CTCGCCCCTGG CTGGCCACCT ACTGGAGAAA GTTCGCATGA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT ATATCCCTCA 'CAGGTTCCTA TCGAACTATGA CCACCATCGA ATATCTATAT TCAACTATCT GAAGAGTCAT TCAACTATCAT TCAACTATCT AAGCCCGGA CCGGAAAGCCT GCCTGAGCGA ATGGTTTCCA ATGGCCATGG GCCATGGCCA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA TCGGATGCAT CGTGCATGAT GTACGGCAT GGACAGCAG ATTATTCTGG GTTCACTTTC CATACGCCT TTCCTCTGCA CTCACATCTG CCTCACATCTG CCTCACATCTG CCTCACATCTG CCTCTGTGCA TTCCTCTACAGC TTCCTCTACAGC TTCCTCTACAGC TTCCCCTCC GGAAGACGGA AAGCCTCGT CAGCCCACT CTCCCCTCC TCTCCCCTCC TCTCCCCTCC TCTCCCCTCC TCTCCCCAGT TGGCCCAGT	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCC GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG CGAGGACTC TCTTTGACGT ACTTCACTCT TCTTTGACGT ACTTCACTCT ACCTCAGACC GACCATGAGA GGGAAGCCC GACCATGAGA GGGAAGCCCA	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCAATTACT CCACTATTAC CCACTATTAC CAATTGCTT TCCTCGTC GGCGCCAGCC AAGTGGTGGT TCCCCATCAC CCACCCCACC	GGGCAACTAC AGGCCACC TTTGGACAGG TGTGCCTGTG GGACTACTT GTTCCCTG GTTTCAGCTT TTGTTCCCTG GAGCTACTT TTGTTCAGCTT TTGTTCAGTT CAAGGAGGTG GAGGGAAGTT CTCTTGGATA AACTTCAATG TTCCTGTATC CTTGCTGAG CCAGCGAGC AGGAAACGTG CCAGCCCAG CCAGCCCAG CCAGCCCAG CCAGCCAG	120 180 240 300 360 420 480 540 600 660 720 840 900 1020 1080 1140 1200 1320 1380 1440 1500 1500
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	CCCAGTCCCA AACCTCTTCA AGCAGTGT AGAATATCTA ATCACGATGT CTGAACTTGA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACTACC TCGTTTTGGA CTCATCCTGA AAGGCCATTG TATGTCTACA AGGACACCC CAGGATGGC CTGGCCACCT ACTGCAGAAA GTTCGCAGAAA GTTCGCATTA GTCGAAGACC CTCGAGAAA GTTCGATGAAC CTCGAAGACC CTCGCAGATAAC GTTCGATGAAA	TCCCGAAATT ACTGCAAAA GATACGATGA TTTATGTCAC TTTTTCATCA CCCTGGACTT GAACGGTGCG AATTCCCTAT AGGACATCAT ATATCCCTCA 'CAGGTTCCTA TTGACCATCA CCACCATCGA ATATCTATAT TCAACTATCA TCAACTATCA TCAACTATCA TGAAGGTCAT TGAACTATCA TCAACTATCA TGAACTATCA TGAACTATCA TGAACTATCA TGAAGACCCGGA ACGCAAGCCT ATGGCCATCGA ATGGTTTCCA ATGGCTTCCA ATGGCTAGCA ATGGTTTCCA ATGGCCATGGCA AGCAGGCCT AAGCAGGCTC	CCTGCTGCTC CCACTTCGAG CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA TCGGATGCAT GTACGCATG GTACAGCAT GTACGCAT GTACAGCAT CCTACATCT CATACGCCT CTACATCTC CTACATCT CCTCTGCA CTCACATCT TTCCTCTGCA TTCCCTCT GGAAGACGGA AAGCCTCGGT CAGCCCACT CAGCCACT CAGCCACT CTCCCCTCC GGCTGATGAC TGTTACCCAT TGCCCCAGT TGCCCCAGT TGCCCCAGT TGCCCCAGT TGCCCCAGT TGCCCCAGT TGGCCCAGT TGGCCCAGT TGGCCCCAGT TGGCCCCCAGT TGGCCCCAGT TGGCCCCAGT TGGCCCCAGT TGGCCCCAGT TGGCCCCAGT TGGCCCCAGT TGGCCCCCAGT TGGCCCCAGT TGGCCCCAGT TGGCCCCCAGT TGGCCCCCAGT TGGCCCCAGT TGGCCCCCAGT TGGCCCCAGT TGGCCCCCAGT TGGCCCCAGT TGGCCCCCAGT TGGCCCCCAGT TGGCCCCAGT TGGCCCCAGT TGCCCCCCAGT TGCCCCCAGT TGCCCCCAGT TGCCCCCCAGT TGCCCCCAGT TGCCCCCAGT TGCCCCCAGT TGCCCCCAGT TGCCCCCAGT TGCCCCCAGT TGCCCCCAGT TGCCCCCCAGT TGCCCCCAGT TGCCCCCCAGT TGCCCCCCAGT TGCCCCCAGT TGCCCCCCCCCC	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCA GCCTGCAACC GATGACAACC GATGACAACC GATGACAACC GCTGCAACC GCCAGGGTGA CCGGATAAGC TTGTTCTTG CCAGGACTC CCCTACCAGC GTCAGCTCT TCTTTGACGT ACCTTCAGAGC ACTACTTTTTC GACCATGAAG AGTATTTTTC GACCATGAAGC GAGAAGCCCA GAGAACCACAG GACAACAATG	CTGTGCCCGA TTCAAAAGAT TGAGGTCCCCG AAATGAATAT TAGCATACTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT TCACAGTTCA TCCACATATTAC CCACTATTAC GGCCCAGCC AAGTGGTGT TCCCCAACAT TGTTCCTGTC GCCCACCCCGCA CAGCCCACCCCCGCA CAGCCCACCCCCACACAT TCCCCAACAT TCCCCATCAC CCACTCCGA CAGCCCAGCC	GGGCAACTAC GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGAGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGGT CCACATGACT CCACATGACT CTCTGGATA AACTTCAATG TTCCTGTATC CTTGCTGGAG AGGAACCTG CCAGCGCAG CCAGCCCAG CCAGCCCAG CCAGCCCAG CCGCCCTGGC CAGCTATGGT CCGCAACCGT GAGCTTCACT GAGCTTCACT GAGCTTCACT CCGCAACCGT GGGCAACCGT GGGCAAAC CTGCCTGCC CGGCAACCGT CGGCCAACCGT CGGCCAACCGT CGGCAACCGT CGGCAACCGT CGGCAACCGT CGGCCAACCGT CGCCAACCGT CGCCCCCCCC CGCCCCCCCACCC CACCCCCCCCC CACCCCCCCC	120 180 240 360 420 480 540 660 720 840 900 960 1020 1140 1200 1260 1320 1380 1440 1500 1560
<ul><li>50</li><li>55</li><li>60</li></ul>	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTAGCA CACCCAGATG GATCTGCATA TACACGGTTG TATTTCTACA AACAGCTACC TCGTTTTGGA CTCATTCTACA AGGCCACTTG AAGGCACCC CAGGATGGCC GCCCCCTGG CTGGCCACCT ACTGGAGAAA GTTCGCTTTA GTCGAAGCCC TCGGATGGCC CTGGATGAGC CTGGATGAGC CTCGATGAGAA AGTTCGCTTTA GTCGAAGACC CTCGAAGACC CTCGAAGACC CTCGAAGACC CTCGAAGAAA AGTTCGCTTAA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGACGTGCG AATTCCCTAT AGACATCAT ATATCCCTCA 'CAGGTTCCTA TTGTGCAAGT TGAACTATGA CCACCATCGA ATATCTATAT TCAACTATG GAAGAGTCAT GAAGAGTCAT GAAGAGTCAT GAAGAGTCAT GAAGAGTCAT AGACCCGGA ATGCCCGGA ATGCCATGGA ATGCCATGGA ATGCCATGGA ATGCCATGGA ATGCCATGGA ATGCCATGCA ATGCCATGGCA ATGCCATGGCA ATGCCATGGCA ATGCCATGGCA ATGCCATGGCA ATGCCATGGCA ATGCAGGCTG AATTCAAGTG	CCTGCTGCTC CCACTTCGAG CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA GACTTGGAAA TCGGATGCAT GTACGCAT GTACGCAT GTACGCAT CCTCACTTC CATACGCCT CTACATCTG CTACATCTG CTCTCTGCA TTCCTCTGCA TTCCTCTGCA TTCCTCTGCA CTCACATCT CGAAGACGA AAGCCTACT TCCCCCTCC GGAAGACCGAT TCTCCCCTC GGCTGATGAC TCTCCCCTT TCTCCCCTC GGCTGATGAC TGGCCCAT TGGCCCAT TGGCCCAT TGGCCCAT TGGCCCAT TGGCCCAT TGGCCCAT TGGCCCAT TGGACCTTGAT	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGTG GTGACTGTGG CGACTCACCA CCTGCAACC GATGACAACC GATGACATGT CTGGGAAGGA ATACTGAAGT ACTGTCCTCA CCGCTACCAGC GTCAGCTCTC TCTTTGACGT ACTTCTCTCT TCTTTGACGT ACTTCTCTCT TCTTTTGACGT ACTTCTCTCT TCTTTTGACGT ACTTCTCTCT ACTTCTCTCT GACCATGAGC AGTATTTTTC GACCATGAAG GGAAACAATG AGTACCTGGG AGTACAATG AGTACCTGGG	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATGCGGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT TCCACTATTAC CCACTATTAC CCACTATTAC CAATTGGCTT TCCCCAACAT TGTTCCTGTC GGGGCCAGCC AGTGGTGGT TCCCCATCACA CCACTCCGA CAGCCCGCA CCACCTCCGA CAGCCCGCA CTACTGAT ATTCCAATGA TGCTTCCCAACAT ATTCCAATGA CCACTTCCGA CAGCCCGGCA CCACCTCCGA AGGCCAGCC CCACTTCCGA AGGCCAGCC AGGCCAGCC AGGCCAGCC AGGCCAGCC AGGCCAGCC AGGCCAGCC AGGCCAGCC AGGCCAGCC AGGCCAGCC AGGCCAGCA AGCACAGACGA ACAAGAGCGA GCCTTAATGA	GGGCAACTAC AGTCGTCCTG GGACTACTG GGACTACTG GGACTACTT GTTTCAGCTT GTTTCAGCTT TTGTTCCCTG GAGCTACTT TTGTTCCCTG GAGGTATGAT CCACATGACT CCACATGACT CTACTTGATT CTCTTGGATG TCCTGTATC CTTGGATG TCCTGTATC CTGCTGGAG CCCCGGCCAG CCCCCGGCCAG CCCCTGGCC CAGCTATGGCT CAGCTATGGT CCGCAACCGT GAGCTTGAGC TGGCGCAAGC CTGCCTTGCC TGATGACCT TGATGACC TGATGACCC TGATGACCC TGATGACCC TGATGACCC TGATGACCC TGATGACCC TGATGACCCT TGATGACCC TGATGACCCT TGATGACCC TGATGACCCT TGATGACCCT TGATGACCCT TGATGACCCT TGATGACCCCT TGATGACCCC TGATGACCC TGATGACCCC TGATGACCCC TGATGACCCC TGATGACCC TGATGACC TGCATC TGATGACC TGATGACC TGATGACC TGATGACC TGATGACC TGCATC TGATGACC TGATGACC TGATGACC TGATGACC TGCATC TGATGACC TGCATC TGATGACC TGATGACC TGCATC TGATGACC TGATGACC TGCATC TGATGACC TGATGACC TGATGACC TGATGACC TGATGACC TGATGACC TGATGACC TGATC TGATGACC TGATGACC TGATC TGATGACC TGATGACC TGATGACC TGATGACC	120 180 240 300 360 420 600 660 720 780 840 900 1020 1140 1200 1380 1380 1440 1500 1560 1620 1680
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTGCA TTGAACAGCA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACAGCTACC CGGTTTTGGA CTCATCCTGA AAGGCCATTG TATGTCTACA AGGAGCCCC CAGGATGGC CTGGCCACCT ACTGGAGAAC GTCGCTTTA GTCGATGGCC TCGGATGGCC TCGGATGGCC TCGGATGAGCC ATTAGAGAGC ATTAGAGAGC ATTAGAGAGC ATGGCCCATG	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AGGACTACTA TCAGCTTCCTA TTGTGCAAGT TCAACTATCA CCACCATCGA ATATCTATAT TCAACTATCT GAAGAGTCAT TCAACTATCA GAAGAGTCAT CAAGCCCGGA CCGAAAGCCT GCCTGAGCGA ATGCCTGACGA ATGCCATGCA ATGCCATGCA ATGCCATGCA ATGCCATGCA ATGCCATGCA ATGCCATGCA ATGCCATGCA AATTCAAGTG GCCATGGCTA AATTCAAGTG GCCAAGAGAA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA AGACTTGAAA TCGGATGCAT CGTGCATGAT GTACGGCAT GTACGGCAT GTACTGCAT GTACAGCAT CTTCACTTCC CTACTGGC TTCCTCTGCA TTCTACAGCT TTCTACAGT TGCCCGCTAC CCAAAGCAG ATTATTCTGG CTTCTACAGT TCCTCTGCA TTCTACAGT TGCCCGCTAC GGAAGACGGA AAGCCTCGGT CAGCCCACTC TCTCCCCCC GGCTGATGAC TGTTACCCAT TGGCCCCAGT TGGCCCCAGT TGGCCCCAGT TGGCCCCAGT TGGCCCCAGT TGACACTTAAC GGACCTGATGAC GGACCTTGAT	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATCTCAGCT GAGAAGTTGT GTGACTGTGG CGACTCACCC GATGACAACC GATGACAACC GATGACAACC GATGACAACA GCCAGGGTGA ACTGTCCTCA GCCAGGGTGA CGGGATAAGC TTCTTCTTCT CGCTACCACC GTCAGCTCT ACCTCAGAGC AGTATTTTC ACCTCAGAGC AGTATTTTC GACCATGAAG GGGAAGCCCA GACAACAATG GACAACAATG AGTACCTGGG TCAGGGTCT TCATGGGG AGTATTTTC GACCATGAAG GGGAAGCCCA GACAACAATG AGTACCTGGG TCAGAGTCTG TCAGAGTCTC	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCAATTACTA CCACTATTAC CCAATTGCTT TCCCCAACAT TGTTCCTGTC GGCGCCAGCC AAGTGGTGGT CCACCTCCGA CCACCTCCGA AGGCCAGCC AGGCCAGCC AGGCCAGCC AGGCCAGCC AGGCCAGCC AAGTGGTGT CCCACTCCGA ATTCCAATGA TGCTTCACCA ATTCCAATGA TGCTTCACCA ACAAGAGCGA ACAAGAGCGA AGGATAGTTG AGGATAGTTG	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGTGCCACT TTTGGACACC CTGCTACTTT GTTTCACGTT TTGTTCCCTG GAGCTATGAT CCACATGACT CAAGGAGGTG GAGGGAAGTT CTCTTGGATA AACTTCAATG TTCCTGTAC CTTGCTGCAC AGGAAACCTG CCCAGCCAG CCCCTGGCC CAGCTATGGT CCGCACCAGC CCGCTTGAC CCGCACCAGC CCGCTTGAC TGGCTACGT TGGCTTGAC TGGCGAACAC TGGCGAACAC TGGCGAACAC TGGCGAACAC TGGCGAACAC TGGCGAACAC TGGCCTTGAC TGGCGAGAAG CTGCCTTGAC TGGCGAGAAG CTGCCTTGAC TGATGAGCTC CCCCCAACCC TGATGAGCTC CCCCCCAACCC CAGCCCAACC	120 180 240 300 360 420 540 600 660 720 780 840 900 1020 11200 1260 1320 1380 1440 1500 1560 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTGCA TTGAACAGCA CACCCAGATG GATCTGCATA TACACGGTTG GATCTGCATA AACAGCTACC TCGTTTTGCA AACAGCTACC TCGTTTTGCA AAGGCCATTG TATGTCTACA AGGAGCCCC CAGGATGGC CTCGCCCTGG CTCGCCCCTGG CTGGCCACCT ACTGGAAGCCC TCGGATGAGC TCGGATGAGC TCGGATGAGC CTCGGATGAGC CCTGGCCCATG CCTGGCCCATG	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT ATATCCCTCA 'CAGGTTCCTA TCGAACTATGA CCACCATCGA ATATCTATAT TCAACTATCT GAAGAGTCAT TGAACTATCT GAAGAGTCAT TGAACTATCT GAAGAGTCAT TGAACTATCT GAAGAGTCAT GAATTAACGT CAAGCCCGGA ACGCATGGCA ATGGTCTCCA ACCAGGGA ACCATCGG GCCATGGCCA AAGCAGGCTG AATTCAACTG AATTCAACTG GCCAAGAGAA CCTTCACTGA	CCTGCTGCTC CCCACTTCGAG TTGTGCAAAT CCGCCTGAGA AGGCATTGAA TCGGATGCAT CGTGCATGAT GTACGGCAT CGTACAGCAT GTACAGCAT CTACAGCAT CTACAGCT TTCCTCTGCA CTCACATCTG CCTCACATCTG CCTCACATCTG CCTCACATCTG CCTCTGTACAG TTTCTACAGCT TTCCTCTACAG TTTCTACAGT TGCCCGCTAC GGAAGACGGA AAGCCTCGT TCTCCCCTCC GGCTGATGAC TCTTCCCCTCC GGCTGATGAC TGTTACCCAT TGGCCCAGT GGACCTCGAT TGGCCCCAGT GGACCTTGAT TGATACTAAC AGGGTTCTCC	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG GCACTCACCA GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG CGAGGACTCTC TCTTTGACGT ACTTCAGCTCT ACTTCAGACT ACTTCAGACT GACCATGAGA GGAAGACCCA GACAACAATG AGTACCTGGG TCAGACTCTC GACCATGAGA GTAACTGAGC TACTCTCTT ACTTCAGACT ACTTCAGACT TTTTTCTCTCT ACCTCAGAGC AGTATTTTTC GACCATGAAG TCAGAGTCTG TCAGATCTCT TTCGATCTCT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCAATTACC CCACTATTAC CCACTATTAC CCACTATTAC GGCCCACCA AGGGCCAGC AAGTGGTGGT TCCCCATCAG CCACCTCCGA CCACCTCCGA AGGCCAGC AGGCCAGCC AAGTGGTGT TCCCATCAC AAGAGAGAA TTCTTCACTA ACTTCACCA ACAAGAGCGA GCCTTCACTA ACAAGAGCGA AGCCTTAATGA TGCTTCACTA ACAAGAGTTG TTAATCCTGA	GGGCAACTAC AGTCCTCTG TTTGGACAGG TGTGCCTGTG GGACTACTTT GTTCACCTT GTTTCAGCTT TTGTTCCCTG GAGCTATGAT CAAGGAGGTG CACATGACT CTCTTTGGATAG AACTTCAATG TTCTTCAGTAT AACTTCAATG TTCCTGTATC CTTGCTGAGA GCAGCCAG GCAGCCAG CCCCCTGGCC CAGCTATGGT GAGGCACAC TGGCTAGGC TGGCGAAC TGGCGAAC TGGCGAAAC TGGCCAAC CTGCCTTGCC CAGCTTGCC CACCTTGCC CCCCCAACC CTACTCCCA	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020 1140 1200 1380 1440 1500 1500 1500 1620 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	CCCAGTCCCA AACCTCTTCA AGCTCTCAA AGAATATCTA ATCACGATGT CTGAACTTGAA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACCTACC TCGTTTTGGA CTCATCCTGA AAGGCCATTG TATTCTACA AGGACACCC CAGGATGGC CTGCACCT ACTGCAGAAA GTTCGCATGA GTTCGATGAGC CTCGATGAGCC CTCGATGAGCC CTCGATGAGC CTCGGATGAGC ATTAAGGAGC ATGGCCCATG	TCCCGAAATT ACTGCAAAAA GATACGATGA TTTATGTCAC TTTTTCATCA CCCTGGACTT GAACGGTGCG AATTCCCTAT ATATCCCTCA 'CAGGTTCCTA TTGACATCAT ATATCCTCA 'CAGGTTCCTA TTGAACTATCA TCAACTATC TCAACTATC TCAACTATC TCAACTATC TCAACTATC TCAACTATC TCATTAACGT CAAGGCCCGGA ATGGTTCCA ATGGCCATCGA ATGGTTTCCA ATGGCCATCGA ATGGTTTCCA ATGGCCATGGA ATGGTTTCCA ATGGCCATGGCA ATGGTTTCCA ATGGCCATGGCA ATGGTTTCCA ATGCCATGGA CCCTCAAGAGAA ACCTTCACTGA AGTGGTCCCC	CCTGCTGCTC CCACTTCGAG CCACTTCGAG AGCATTGAAA TCGGCTGAGAA TCGGATGCAT GTACGCATGAT GTACGCATGAT GTACGCATGCAT GTACACACTC CTACATCTC CTACATCTC CTACATCTGC CTACATCTGC CTCCTGTGTG TTCCTCTGCA CTCACATCT CTCCCCTCC GGAAGACGGA AGCCTCGCT CAGCCCCC CAGCTC CAGCCCCCC GGCTGATGAC TGTCCCCTTC GGCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGACCTTGAT TGATACTAAC GGACATTACC GGACATTACC GGACATTACC GGACATTACC GGACATTACC GGACCTTCCTCC GGTTCTCCCCTCC GGTTCTCCCCTCC GGTTCTCCCCTCC GGACCTTGAT TGATACTAAC GGACAGTAGC GGTTCCTCTC	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG GCACTCACCA GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG CGAGGACTCTC TCTTTGACGT ACTTCAGCTCT ACTTCAGACT ACTTCAGACT GACCATGAGA GGAAGACCCA GACAACAATG AGTACCTGGG TCAGACTCTC GACCATGAGA GTAACTGAGC TACTCTCTT ACTTCAGACT ACTTCAGACT TTTTTCTCTCT ACCTCAGAGC AGTATTTTTC GACCATGAAG TCAGAGTCTG TCAGATCTCT TTCGATCTCT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCAATTACC CCACTATTAC CCACTATTAC CCACTATTAC GGCCCACCA AGGGCCAGC AAGTGGTGGT TCCCCATCAG CCACCTCCGA CCACCTCCGA AGGCCAGC AGGCCAGCC AAGTGGTGT TCCCATCAC AAGAGAGAA TTCTTCACTA ACTTCACCA ACAAGAGCGA GCCTTCACTA ACAAGAGCGA AGCCTTAATGA TGCTTCACTA ACAAGAGTTG TTAATCCTGA	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGTGCCACT TTTGGACACC CTGCTACTTT GTTTCACGTT TTGTTCCCTG GAGCTATGAT CCACATGACT CAAGGAGGTG GAGGGAAGTT CTCTTGGATA AACTTCAATG TTCCTGTAC CTTGCTGCAC AGGAAACCTG CCCAGCCAG CCCCTGGCC CAGCTATGGT CCGCACCAGC CCGCTTGAC CCGCACCAGC CCGCTTGAC TGGCTACGT TGGCTTGAC TGGCGAACAC TGGCGAACAC TGGCGAACAC TGGCGAACAC TGGCGAACAC TGGCGAACAC TGGCCTTGAC TGGCGAGAAG CTGCCTTGAC TGGCGAGAAG CTGCCTTGAC TGATGAGCTC CCCCCAACCC TGATGAGCTC CCCCCCAACCC CAGCCCAACC	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020 1140 1200 1380 1440 1500 1500 1500 1620 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	CCCAGTCCCA AACCTCTTCA AGCTCTCAA AGAATATCTA ATCACGATGT CTGAACTTGAA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACCTACC TCGTTTTGGA CTCATCCTGA AAGGCCATTG TATTCTACA AGGACACCC CAGGATGGC CTGCACCT ACTGCAGAAA GTTCGCATGA GTTCGATGAGC CTCGATGAGCC CTCGATGAGCC CTCGATGAGC CTCGGATGAGC ATTAAGGAGC ATGGCCCATG	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT ATATCCCTCA 'CAGGTTCCTA TCGAACTATGA CCACCATCGA ATATCTATAT TCAACTATCT GAAGAGTCAT TGAACTATCT GAAGAGTCAT TGAACTATCT GAAGAGTCAT TGAACTATCT GAAGAGTCAT GAATTAACGT CAAGCCCGGA ACGCATGGCA ATGGTCTCCA ACCAGGGA ACCATCGG GCCATGGCCA AAGCAGGCTG AATTCAACTG AATTCAACTG GCCAAGAGAA CCTTCACTGA	CCTGCTGCTC CCACTTCGAG CCACTTCGAG AGCATTGAAA TCGGCTGAGAA TCGGATGCAT GTACGCATGAT GTACGCATGAT GTACGCATGCAT GTACACACTC CTACATCTC CTACATCTC CTACATCTGC CTACATCTGC CTCCTGTGTG TTCCTCTGCA CTCACATCT CTCCCCTCC GGAAGACGGA AGCCTCGCT CAGCCCCC CAGCTC CAGCCCCCC GGCTGATGAC TGTCCCCTTC GGCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGCCCCAT TGGACCTTGAT TGATACTAAC GGACATTACC GGACATTACC GGACATTACC GGACATTACC GGACATTACC GGACCTTCCTCC GGTTCTCCCCTCC GGTTCTCCCCTCC GGTTCTCCCCTCC GGACCTTGAT TGATACTAAC GGACAGTAGC GGTTCCTCTC	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG GCACTCACCA GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG CGAGGACTCTC TCTTTGACGT ACTTCAGCTCT ACTTCAGACT ACTTCAGACT GACCATGAGA GGAAGACCCA GACAACAATG AGTACCTGGG TCAGACTCTC GACCATGAGA GTAACTGAGC TACTCTCTT ACTTCAGACT ACTTCAGACT TTTTTCTCTCT ACCTCAGAGC AGTATTTTTC GACCATGAAG TCAGAGTCTG TCAGATCTCT TTCGATCTCT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCAATTACC CCACTATTAC CCACTATTAC CCACTATTAC GGCCCACCA AGGGCCAGC AAGTGGTGGT TCCCCATCAG CCACCTCCGA CCACCTCCGA AGGCCAGC AGGCCAGCC AAGTGGTGT TCCCATCAC AAGAGAGAA TTCTTCACTA ACTTCACCA ACAAGAGCGA GCCTTCACTA ACAAGAGCGA AGCCTTAATGA TGCTTCACTA ACAAGAGTTG TTAATCCTGA	GGGCAACTAC AGTCCTCTG TTTGGACAGG TGTGCCTGTG GGACTACTTT GTTCACCTT GTTTCAGCTT TTGTTCCCTG GAGCTATGAT CAAGGAGGTG CACATGACT CTCTTTGGATAG AACTTCAATG TTCTTCAGTAT AACTTCAATG TTCCTGTATC CTTGCTGAGA GCAGCCAG GCAGCCAG CCCCCTGGCC CAGCTATGGT GAGGCACAC TGGCTAGGC TGGCGAAC TGGCGAAC TGGCGAAAC TGGCCAAC CTGCCTTGCC CAGCTTGCC CACCTTGCC CCCCCAACC CTACTCCCA	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020 1140 1200 1380 1440 1500 1500 1500 1620 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li></ul>	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTGCA TTGAACAGCA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACAGCTACC CCGTTTTGGA AAGGCCATTG TATGTCTACA AGGAGCCCC CAGGATGGC CTGGCCACCT ACTGGAGAAC GTCCCCTTGG GTCGCAAGCC TCGGATGGCC TCGGATGGCC TCGGATGAGA ATTAAGAGC ATTAAGAGA ATTAGGACC CTGGGTGCCATG CTGGCCATG CTGGCCATG CTGGCTGAAC ATTAGGAGC ATTAGGAGC ATGGCCCATG CCTGGGTGCCATG CCTGGGTGCCATG CCTGGGTGCCATG CCTGGGTGCCATG CCTGGGTGCCATG CCTGGGTGCCATG CCTGGGTGCCATG CCTGGGTGCCA AAGGTCGACA TACTGGGTACA TACTGGGTACA	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AGGACTACTA TCAGCTTCCTA TTGTGCAAGT TCAACTATCA TCAACTATCA TCAACTATCT GAAGAGTCAT TCAACTATCA GAAGAGTCAT TCAACTATCA GAAGAGTCAT AGGCCCGGA ACTGCCAAGAGAA ATGCCTTAACGT GCCATGGCCA ATGCCATGGA ATGCCATGCA ATGCCATGCA ATGCCATGCA ATGCCATGCA ACCATGACAAC CCTTCACTGA ACTGCTCACTGA ACTGCTCACTGA ACTGCTCACTGA ACTGCTCACTGA ACTGCTCCCGA ACCATATGTA	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA AGACTTGAAA TCGGATGCAT CGTGCATGAT GTACGGCAT GTACGGCAT GTACTGCAT GTACAGCAT CATACGCCT CTACTGGA ATTATTCTGG GTTCACTTTC CATACGCCT TTCCTCTGCA TTCCTCTGCA TTCTACAGT TGCCCGCTAC CGAAAGACGG AAGCCTCGGTAC CGAAGCAGG CAGCCCACTC TCTCCCCCC GGCTGATGAC TGTTACCCAT TGGCCCAGT GGACCTTGAT TGGCCCCAGT GGACCTTGAT TGATACTAC GGAAGATGAC GGAACGTAAC GGACAGTAGC GGACGTAC GGACGTATCC GTTCCTCCC GTTCCTCTC TTAG	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG GCACTCACCA GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG CGAGGACTCTC TCTTTGACGT ACTTCAGCTCT ACTTCAGACT ACTTCAGACT GACCATGAGA GGAAGACCCA GACAACAATG AGTACCTGGG TCAGACTCTC GACCATGAGA GTAACTGAGC TACTCTCTT ACTTCAGACT ACTTCAGACT TTTTTCTCTCT ACCTCAGAGC AGTATTTTTC GACCATGAAG TCAGAGTCTG TCAGATCTCT TTCGATCTCT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCAATTACC CCACTATTAC CCACTATTAC CCACTATTAC GGCCCACCA AGGGCCAGC AAGTGGTGGT TCCCCATCAG CCACCTCCGA CCACCTCCGA AGGCCAGC AGGCCAGCC AAGTGGTGT TCCCATCAC AAGAGAGAA TTCTTCACTA ACTTCACCA ACAAGAGCGA GCCTTCACTA ACAAGAGCGA AGCCTTAATGA TGCTTCACTA ACAAGAGTTG TTAATCCTGA	GGGCAACTAC AGTCCTCTG TTTGGACAGG TGTGCCTGTG GGACTACTTT GTTCACCTT GTTTCAGCTT TTGTTCCCTG GAGCTATGAT CAAGGAGGTG CACATGACT CTCTTTGGATAG AACTTCAATG TTCTTCAGTAT AACTTCAATG TTCCTGTATC CTTGCTGAGA GCAGCCAG GCAGCCAG CCCCCTGGCC CAGCTATGGT GAGGCACAC TGGCTAGGC TGGCGAAC TGGCGAAC TGGCGAAAC TGGCCAAC CTGCCTTGCC CAGCTTGCC CACCTTGCC CCCCCAACC CTACTCCCA	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020 1140 1200 1380 1440 1500 1500 1500 1620 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	CCCAGTCCCA AACCTCTTCA AGCTCTCAA AGAATATCTA ATCAGGATGT CTGAACTTGG CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACTACC TCGTTTTGGA CTCATCCTGA AAGGCCATTG TATTCTACA AGGACACCC CAGGATGGCC CAGGATGGCC CTGGTTTACA AGTCCCTTGG CTCGCACCT ACTGCAGAAA GTTCGCAGACC CTCGGATGAGC CTCGCATGGGCACCT ACTGCAGAGAA ATTAAGGAGC CTCGCATGGCCATGGCACAT ACTGCCATGC ATGCCCATG	TCCCGAAATT ACTGCAAAAA GATACGATGA TTTATGTCAC TTTTTCATCA CCCTGGACTT GAACGGTGCG AATTCCCTAT ATGACATCAT ATGACATCAT ATATCCCTCA 'CAGGTTCCTA TTGACAAGT TCAACTATCA CCACCATCGA ATATCTATAT TCAACTATCT GAAGAGTCAT TGATTAACGT CAAGCCCGGA ATGATTACCT ATGACCATCAA ATGATTACA TGATTAACGT CAAGCCCGGA ATGGTTTCCA ATGGCCATGG AATGACAGCA ATGGTTTCCA AAGCAGGCTG AATTCAAGTG GCCAAGAGGA ACCATCAGCA AGCAGGCTG AATTCAAGTG ACCATCACTGA AGTGGTCCCG ACCATATGTA 38 Protein	CCTGCTGCTC CCCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA TCGGATGCAT GTACGGCAT GTACGGCAT GTACAGCAT GTACAGCAT CCTCACATCTG CTACATCTG CTCACATCTG CTCACATCTG CTCACATCTG CCTCGTGTGC TTCCTCTACAG TTCCCCTCG GAAGACGGA AAGCCTCGGT CAGCCCACT TCTCCCCTC GGCTGATGAC TGTTACCCAT TGGCCCACT TGTACCAT TGGCCCCAGT GGACCTCAT GGACCATCAGAT TGATACTAAC GGACAGTAGC GGACCTCCC GTTCCTCTC TTAG  Eequence	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG GCACTCACCA GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG CGAGGACTCTC TCTTTGACGT ACTTCAGCTCT ACTTCAGACT ACTTCAGACT GACCATGAGA GGAAGACCCA GACAACAATG AGTACCTGGG TCAGACTCTC GACCATGAGA GTAACTGAGC TACTCTCTT ACTTCAGACT ACTTCAGACT TTTTTCTCTCT ACCTCAGAGC AGTATTTTTC GACCATGAAG TCAGAGTCTG TCAGATCTCT TTCGATCTCT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCAATTACC CCACTATTAC CCACTATTAC CCACTATTAC GGCCCACCA AGGGCCAGC AAGTGGTGGT TCCCCATCAG CCACCTCCGA CCACCTCCGA AGGCCAGC AGGCCAGCC AAGTGGTGT TCCCATCAC AAGAGAGAA TTCTTCACTA ACTTCACCA ACAAGAGCGA GCCTTCACTA ACAAGAGCGA AGCCTTAATGA TGCTTCACTA ACAAGAGTTG TTAATCCTGA	GGGCAACTAC AGTCCTCTG TTTGGACAGG TGTGCCTGTG GGACTACTTT GTTCACCTT GTTTCAGCTT TTGTTCCCTG GAGCTATGAT CAAGGAGGTG CACATGACT CTCTTTGGATAG AACTTCAATG TTCTTCAGTAT AACTTCAATG TTCCTGTATC CTTGCTGAGA GCAGCCAG GCAGCCAG CCCCCTGGCC CAGCTATGGT GAGGCACAC TGGCTAGGC TGGCGAAC TGGCGAAC TGGCGAAAC TGGCCAAC CTGCCTTGCC CAGCTTGCC CACCTTGCC CCCCCAACC CTACTCCCA	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020 1140 1200 1380 1440 1500 1500 1500 1620 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	CCCAGTCCCA AACCTCTTCA AGCTCTCAA AGAATATCTA ATCAGGATGT CTGAACTTGG CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACTACC TCGTTTTGGA CTCATCCTGA AAGGCCATTG TATTCTACA AGGACACCC CAGGATGGCC CAGGATGGCC CTGGTTTACA AGTCCCTTGG CTCGCACCT ACTGCAGAAA GTTCGCAGACC CTCGGATGAGC CTCGCATGGGCACCT ACTGCAGAGAA ATTAAGGAGC CTCGCATGGCCATGGCACAT ACTGCCATGC ATGCCCATG	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AGGACTACTA TCAGCTTCCTA TTGTGCAAGT TCAACTATCA TCAACTATCA TCAACTATCT GAAGAGTCAT TCAACTATCA GAAGAGTCAT TCAACTATCA GAAGAGTCAT AGGCCCGGA ACTGCCAAGAGAA ATGCCTTAACGT GCCATGGCCA ATGCCATGGA ATGCCATGCA ATGCCATGCA ATGCCATGCA ATGCCATGCA ACCATGACAAC CCTTCACTGA ACTGCTCACTGA ACTGCTCACTGA ACTGCTCACTGA ACTGCTCACTGA ACTGCTCCCGA ACCATATGTA	CCTGCTGCTC CCCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA TCGGATGCAT GTACGGCAT GTACGGCAT GTACAGCAT GTACAGCAT CCTCACATCTG CTACATCTG CTCACATCTG CTCACATCTG CTCACATCTG CCTCGTGTGC TTCCTCTACAG TTCCCCTCG GAAGACGGA AAGCCTCGGT CAGCCCACT TCTCCCCTC GGCTGATGAC TGTTACCCAT TGGCCCACT TGTACCAT TGGCCCCAGT GGACCTCAT GGACCATCAGAT TGATACTAAC GGACAGTAGC GGACCTCCC GTTCCTCTC TTAG  Eequence	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG GCACTCACCA GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG CGAGGACTCTC TCTTTGACGT ACTTCAGCTCT ACTTCAGACT ACTTCAGACT GACCATGAGA GGAAGACCCA GACAACAATG AGTACCTGGG TCAGACTCTC GACCATGAGA GTAACTGAGC TACTCTCTT ACTTCAGACT ACTTCAGACT TTTTTCTCTCT ACCTCAGAGC AGTATTTTTC GACCATGAAG TCAGAGTCTG TCAGATCTCT TTCGATCTCT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCAATTACC CCACTATTAC CCACTATTAC CCACTATTAC GGCCCACCA AGGGCCAGC AAGTGGTGGT TCCCCATCAG CCACCTCCGA CCACCTCCGA AGGCCAGC AGGCCAGCC AAGTGGTGT TCCCATCAC AAGAGAGAA TTCTTCACTA ACTTCACCA ACAAGAGCGA GCCTTCACTA ACAAGAGCGA AGCCTTAATGA TGCTTCACTA ACAAGAGTTG TTAATCCTGA	GGGCAACTAC AGTCCTCTG TTTGGACAGG TGTGCCTGTG GGACTACTTT GTTCACCTT GTTTCAGCTT TTGTTCCCTG GAGCTATGAT CAAGGAGGTG CACATGACT CTCTTTGGATAG AACTTCAATG TTCTTCAGTAT AACTTCAATG TTCCTGTATC CTTGCTGAGA GCAGCCAG GCAGCCAG CCCCCTGGCC CAGCTATGGT GAGGCACAC TGGCTAGGC TGGCGAAC TGGCGAAC TGGCGAAAC TGGCCAAC CTGCCTTGCC CAGCTTGCC CACCTTGCC CCCCCAACC CTACTCCCA	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020 1140 1200 1380 1440 1500 1500 1500 1620 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	CCCAGTCCCA AACCTCTTCA AGCTCTCAA AGAATATCTA ATCAGGATGT CTGAACTTGG CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACTACC TCGTTTTGGA CTCATCCTGA AAGGCCATTG TATTCTACA AGGACACCC CAGGATGGCC CAGGATGGCC CTGGTTTACA AGTCCCTTGG CTCGCACCT ACTGCAGAAA GTTCGCAGACC CTCGGATGAGC CTCGCATGGGCACCT ACTGCAGAGAA ATTAAGGAGC CTCGCATGGCCATGGCACAT ACTGCCATGC ATGCCCATG	TCCCGAAATT ACTGCAAAAA GATACGATGA TTTATGTCAC TTTTTCATCA CCCTGGACTT GAACGGTGCG AATTCCCTAT ATGACATCAT ATGACATCAT ATATCCCTCA 'CAGGTTCCTA TTGACAAGT TCAACTATCA CCACCATCGA ATATCTATAT TCAACTATCT GAAGAGTCAT TGATTAACGT CAAGCCCGGA ATGATTACCT ATGACCATCAA ATGATTACA TGATTAACGT CAAGCCCGGA ATGGTTTCCA ATGGCCATGG AATGACAGCA ATGGTTTCCA AAGCAGGCTG AATTCAAGTG GCCAAGAGGA ACCATCAGCA AGCAGGCTG AATTCAAGTG ACCATCACTGA AGTGGTCCCG ACCATATGTA 38 Protein	CCTGCTGCTC CCCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA TCGGATGCAT GTACGGCAT GTACGGCAT GTACAGCAT GTACAGCAT CCTCACATCTG CTACATCTG CTCACATCTG CTCACATCTG CTCACATCTG CCTCGTGTGC TTCCTCTACAG TTCCCCTCG GAAGACGGA AAGCCTCGGT CAGCCCACT TCTCCCCTC GGCTGATGAC TGTTACCCAT TGGCCCACT TGTACCAT TGGCCCCAGT GGACCTCAT GGACCATCAGAT TGATACTAAC GGACAGTAGC GGACCTCCC GTTCCTCTC TTAG  Eequence	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG GCACTCACCA GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG CGAGGACTCTC TCTTTGACGT ACTTCAGCTCT ACTTCAGACT ACTTCAGACT GACCATGAGA GGAAGACCCA GACAACAATG AGTACCTGGG TCAGACTCTC GACCATGAGA GTAACTGAGC TACTCTCTT ACTTCAGACT ACTTCAGACT TTTTTCTCTCT ACCTCAGAGC AGTATTTTTC GACCATGAAG TCAGAGTCTG TCAGATCTCT TTCGATCTCT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCAATTACC CCACTATTAC CCACTATTAC CCACTATTAC GGCCCACCA AGGGCCAGC AAGTGGTGGT TCCCCATCAG CCACCTCCGA CCACCTCCGA AGGCCAGC AGGCCAGCC AAGTGGTGT TCCCATCAC AAGAGAGAA TTCTTCACTA ACTTCACCA ACAAGAGCGA GCCTTCACTA ACAAGAGCGA AGCCTTAATGA TGCTTCACTA ACAAGAGTTG TTAATCCTGA	GGGCAACTAC AGTCCTCTG TTTGGACAGG TGTGCCTGTG GGACTACTTT GTTCACCTT GTTTCAGCTT TTGTTCCCTG GAGCTATGAT CAAGGAGGTG CACATGACT CTCTTTGGATAG AACTTCAATG TTCTTCAGTAT AACTTCAATG TTCCTGTATC CTTGCTGAGA GCAGCCAG GCAGCCAG CCCCCTGGCC CAGCTATGGT GAGGCACAC TGGCTAGGC TGGCGAAC TGGCGAAC TGGCGAAAC TGGCCAAC CTGCCTTGCC CAGCTTGCC CACCTTGCC CCCCCAACC CTACTCCCA	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020 1140 1200 1380 1440 1500 1500 1500 1620 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li><li>75</li></ul>	CCCAGTCCCA AACCTCTTCA AGCTCTCAA AGAATATCTA ATCAGGATGT CTGAACTTGG CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACTACC TCGTTTTGGA CTCATCCTGA AAGGCCATTG TATTCTACA AGGACACCC CAGGATGGCC CAGGATGGCC CTGGTTTACA AGTCGCAGAC GCTCCCTGG CTGCCACCT GCTCGAGAAA GTTCGATGAGC CTCGGATGAGC ATTAAGGAGC ATGCCCATG ATGCCCATG CCTGGGTGCA ATTAGGAGCA TACTGGGTAT Seq ID NO:	TCCCGAAATT ACTGCAAAAA GATACGATGA TTTATGTCAC TTTTTCATCA CCCTGGACTTA AGGATGCTTT GAACGGTGCG AATTCCCTAT ATATCCCTCA 'CAGGTTCCTA TTGTGCAAGT TCAACTATCA ATATCATATA TCAACTATCA CAACCATCGA ATATCTATAT TCAACTATCT GAAGAGTCAT TGATTAACGT CAAGCCCGGA ATGGTTCCA ATGGCCATCGA ATGGTTTCCA ATGGCCATCGA ATGGTTTCCA ATGGCCATCGA ATGGTTTCCA ATGGCATGGC	CCTGCTGCTC CCCACTTCGAG TTGTGCAAAT CCGCCTGAGA GAGCATTGAA TCGGATGCAT GTACGGCAT GTACGGCAT GTACAGCAT GTACAGCAT CCTCACATCTG CTACATCTG CTCACATCTG CTCACATCTG CTCACATCTG CCTCGTGTGC TTCCTCTACAG TTCCCCTCG GAAGACGGA AAGCCTCGGT CAGCCCACT TCTCCCCTC GGCTGATGAC TGTTACCCAT TGGCCCACT TGTACCAT TGGCCCCAGT GGACCTCAT GGACCATCAGAT TGATACTAAC GGACAGTAGC GGACCTCCC GTTCCTCTC TTAG  Eequence	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG GCACTCACCA GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTTG CGAGGACTCTC TCTTTGACGT ACTTCAGCTCT ACTTCAGACT ACTTCAGACT GACCATGAGA GGAAGACCCA GACAACAATG AGTACCTGGG TCAGACTCTC GACCATGAGA GTAACTGAGC TACTCTCTT ACTTCAGACT ACTTCAGACT TTTTTCTCTCT ACCTCAGAGC AGTATTTTTC GACCATGAAG TCAGAGTCTG TCAGATCTCT TTCGATCTCT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT CCAATTACC CCACTATTAC CCACTATTAC CCACTATTAC GGCCCACCA AGGGCCAGC AAGTGGTGGT TCCCCATCAG CCACCTCCGA CCACCTCCGA AGGCCAGC AGGCCAGCC AAGTGGTGT TCCCATCAC AAGAGAGAA TTCTTCACTA ACTTCACCA ACAAGAGCGA GCCTTCACTA ACAAGAGCGA AGCCTTAATGA TGCTTCACTA ACAAGAGTTG TTAATCCTGA	GGGCAACTAC AGTCCTCTG TTTGGACAGG TGTGCCTGTG GGACTACTTT GTTCACCTT GTTTCAGCTT TTGTTCCCTG GAGCTATGAT CAAGGAGGTG CACATGACT CTCTTTGGATAG AACTTCAATG TTCTTCAGTAT AACTTCAATG TTCCTGTATC CTTGCTGAGA GCAGCCAG GCAGCCAG CCCCCTGGCC CAGCTATGGT GAGGCACAC TGGCTAGGC TGGCGAAC TGGCGAAC TGGCGAAAC TGGCCAAC CTGCCTTGCC CAGCTTGCC CACCTTGCC CCCCCAACC CTACTCCCA	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020 1140 1200 1380 1440 1500 1500 1500 1620 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li></ul>	CCCAGTCCCA AACCTCTTCA AGCTCTCAA AGAATATCTA ATCAGGATGT CTGAACTTGG CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACTACC TCGTTTTGGA CTCATCCTGA AAGGCCATTG TATTCTACA AGGACACCC CAGGATGGCC CAGGATGGCC CTGGTTTACA AGTCGCAGAC GCTCCCTGG CTGCCACCT GCTCGAGAAA GTTCGATGAGC CTCGGATGAGC ATTAAGGAGC ATGCCCATG ATGCCCATG CCTGGGTGCA ATTAGGAGCA TACTGGGTAT Seq ID NO:	TCCCGAAATT ACTGCAAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AGGACATCAT TTGTGCAAGT TGAACTATGA CCACCATCGA ATATCTATAT TCAACTATT GAAGAGTCAT GAAGAGTCAT GAAGAGTCAT GAAGAGTCAT GAAGAGTCAT GCCTGAGCGA ATGCCATGGA ATGCCATGGA ATGCCATGCA ATGCCATGCA AGCAGGCTG ACTTCACTGA AGCAGGCTG ACTTCACTGA AGCAGGCTG ACTTCACTGA AGCAGGCTG ACTTCACTGA AGTGGTCCCG ACCATATGTA  38 Protein Cession #: /	CCTGCTGCTC CCACTTCGAG TTGTGCAAAT CCGCCTGAGA AGACTTGAAA TCGGATGCAT GTACGGCAT GTACGGCAT GTACGGCAT GTACGGCAT CCTGCAAG ATTATTCTGG GTTCACTTTC CATACGCCT TTCCTCTGCA TTCACATCT TCCCCTCC GGAAGACGGA AAGCCTGGTAT CGTGATGAC TTTCCCCTCC TCTCCCCTCC TCTCCCCTCC TCTCCCCTCC TCTCCCTCC TTAGGCCCAGT GGACCTGAT GGACCTTGAT TGATACTAAC GGACAGTAGC TTAGC TTAG Sequence AADS1172.1	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATCTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCC GATGACACC GATGACAATG CTGGGAAGGA ATACTGAAGT ACTGTCCTCA GCCAGGGTGA CCGGATAAGC GTCAGCTCTCTTTG CGAGGACCTC GTCAGCTCTC ACCTCAGAGC GTCAGCTCTC ACCTCAGAGC GACAACAATG ACTTCTCTCT GACCATGAAG GGAAGCACA GACAACAATG AGTATTTTC GACCATGAAG GTCAGCTCC GACAACAATG ACTACCTGGG TCAGAGTCTC TCTGATCTCT CCTCTGGCCT	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TGGTGGTAGA GGAACGCCAT TCCACTATTAC CCACTATTAC CCACTATTAC GGGCCAGCC AAGTGGTGGT TCCCCAACAT TCCCATCCGA CAGCCCGCA CCACCTCCGA CAGCCCGGCA CCACCTCCGA AGGCCAGCC CCACTTCCGA AGTGGTGGT TCCCCATCAC CCACTCCGA AGGCCAGCC AGCCTTCCGA ATTCCAATGA ATTCCAATGA ACAAGACGA ACCATAATGA ACAAGACGA ACCTTAATCA TTAATCCTGA TTGGGTTGTT	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGTGCCACT TTTGGACACG TGGGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGGT CCACATGACT CAAGGAGGTG GAGGGAAGTT CTCTTGGATA AACTTCAATG TTCCTGTATC CTTGCTGGAG AGGAACCTG CCCAGCCAG CCCCTGGCC CAGCTATGGC CCGCAACCGT GAGCTTTGAGC TGGCGAGAAG CTGCCTTGCC TGACTGAGC TGAGGAGAAG CTGCCTTGCC TGATGAGCT CCCCAACCCT CAACATTGTT CCCCCAACCCC CAACATTGTT	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020 1140 1200 1380 1440 1500 1500 1500 1620 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li><li>75</li></ul>	CCCAGTCCCA AACCTCTTCA AGCTCTCAA AGAATATCTA ATCACGATGT CTGAACTTGAA CACCCAGATG GATCTGCATA TACACGGTTG GAGGAGCTGC TATTTCTACA AACACTACC TCGTTTTGGA CTCATCCTGA AAGGCCATTG TATGTCTTACA AGGACACCC CAGGATGGC CTGGCCACCT ACTGCAGAAA GTTCGATGAAC GTTCGATGAGC CTCGATGAGC CTCGGATGAGC CTCGGATGAGC ATTAAGGAGC ATTAAGGACC TCGGATGAGC ATTAAGGACC TCGGATGAGC ATTAAGGACC TCGGATGAGC ATTAAGGACC TCGGATGAGC ATTAAGGACC TCGGATGAGC ATTAAGGACC TCGGATGAGC TCCTGGGTTCCAAG ATTAAGGACC TCGGATGACA ATTAAGGACC TCCTGGGTTCCAAG TCCTGGTTCCAAG TCCTGGTTCCAAG TCCTGGTTCCAAG TCCTGGTTCCAAG TCCTGGTTCCAAG TCCTGGTTCCAAG TCCTGGTTCCAAG TCCTGTTCCAAG TCCTGGTTCCAAG TCCTGTTCCAAG TCCTCTCTGAAG TCCTCTGTTCCAAG TCCTGTTCCAAG TCCTGTTCCAAG TCCTGTTCCAAG TCCTGTTCCAAG TCCTGTTCCAAG TC	TCCCGAAATT ACTGCAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTT GAACGGTGCG AATTCCCTAT ATGACATCAT ATGACATCAT ATGACATCAT TCGACATCAT CCACCATCGA ATTATCATTAT TCAACTATC TGAAGGTCAT TGAACTATC TGAAGACTAT TCAACTATC TGAAGACTAT TCAACTATC TGAAGACCCGGA ATGATTACCT ATTGCCATGGA ATGGTTTCCA ATGGCATGGA ATGGTTTCCA ATGGCATGGA ATGGTTTCCA AGCGCTG AATTCAAGTG GCCAAGAGGA ACCTTCACTGA AGTGGTCCCG ACCATATGTA  38 Protein cession #: 1	CCTGCTGCTC CCCACTTCGAG TTGTGCAAAT CCGCCTGAGA AGCATTGAAA TCGGATGCAT GTACGGCAT GTACGGCAT GTACAGCAT CCTCACATCTG CTACATCTGC CTCACATCTG CTCACATCTG CCTCGTGTGC TTCCTCTACAG TTTCCCCTCG GCAAGACGGA AAGCCTCGGT TCCCCTCC GGCTGATGAC TCTCCCCTCC GGCTGATGAC TCTCCCCTCC GGCTGATGAC TGTTACCCAT TCACATTTC TCTCCCCTCC GGCTGATGAC TGTTACCCAT TGACCCACT TTTACCCAT TGACCCCAGT GGACCTTGAT TGATACTAAC GGACAGTAGC AGGATTCTC TTAG  Sequence AADS1172.1	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCA GCCTGCAACC GATGACACC GATGACACT GAGACATGTGA ACTGTCCTCA GCCAGGGTGA CCGGATAAGC TTGTTCTTCA CCTCAGCTCT TCTTTGACGT ACTTCTCTCT ACCTCAGAGC AGTATTTTC CGACCATGAGG GACAACAATG ACTACCTGGG TCAGCTCTC GACCATGAGG TCAGCTCTC CCTCTGGCCT CCTCTGGCCT CCTCTGGCCT CCTCTGGCCT CCTCTGGCCT CCTCTGGCCT	CTGTGCCCGA TTCAAAAGAT TTCAAAAGAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC CTACAGCAGC TGGTGGTAGA GGAACGCCAT TCCCCAACAT TCCCCAACAT TCCCCAACAT TCCCCATCAGC CAATTGGGTG AGGACGCCAT CCACTCCGA ACTGGTGC CAACTCCGA ACTGGTGCA CAACTCCGAACT TCCCCATCAC CAACTCCGAACT TCCCCATCAC CAACTCCGAACT TCCCCATCAC CAACTCCGAACT TCCCCATCAC CAACTCCGAACT TCCCCATCAC CAACTCCGAACT TTCCAATGA AGGCCAGGC AGCCCAGCC ACAAGAGCGA TTTCCATCAC TCTACCGAAT TTCCATCAC TTGCGTTGTT  41	GGGCAACTAC AGTCGTCCTG TTTGGACAGG TGTGCCTGTG GGACTACACG TGTGCCACT TTTGGACACG TGGGACCACC CTGCTACTTT GTTTCAGCTT TTGTTCCCTG GAGCTATGGT CCACATGACT CAAGGAGGTG GAGGGAAGTT CTCTTGGATA AACTTCAATG TTCCTGTATC CTTGCTGGAG AGGAACCTG CCCAGCCAG CCCCTGGCC CAGCTATGGC CCGCAACCGT GAGCTTTGAGC TGGCGAGAAG CTGCCTTGCC TGACTGAGC TGAGGAGAAG CTGCCTTGCC TGATGAGCT CCCCAACCCT CAACATTGTT CCCCCAACCCC CAACATTGTT	120 180 240 300 360 420 480 540 600 660 720 840 900 900 1020 1140 1200 1380 1440 1500 1500 1500 1620 1680 1740
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li><li>75</li></ul>	CCCAGTCCCA AACCTCTTCA AGCTCTCAA AGAATATCTA ATCACGATGT CTGAACTIGA CACCCAGATG GATCTGCATA TACACGGTTG TATCTACA AACAGCTAC TCGTTTTGGA CTCATCTGGA ACTCCTGGA AAGGCCATTG TATGTCTACA AGGGCACTG CTGGCCACCT CGGTGGCCACCT TCGTTGGA GTTCGCATGGA ATTAAGGAGA ATTAAGGAGA ATTAAGGAGC CTCGGTTGCACAC TCGGTTGCAC TCGTGTGAGAC CTCGATGAGC TCGCATGAGC TCGATGAGC TCGCATGAGC TCGCTGCT TAAGGTCGAAC ATTAAGGAGC TCGCTGGTTGC TCGGTTGCAAC TACTGGGTGAC TACTGGGTGAC TACTGGGTAT  Seq ID NO: Protein Ac  I   MLRAAVILLL MLRAAVILLL	TCCCGAAATT ACTGCAAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTT GAACGGTGCG AATTCCCTAT AGGACATCAT ATTCCCTCA 'CAGGTTCCTA TTGTGCAAGT TGAACTATTA TCAACTATT GAAGAGTCAT TCAACTATT GAAGAGTCAT TCAACTATCT GAAGAGTCAT CAAGCCCGGA ATATCTATAT TCAATTAACGT CAAGCCCGGA ATGCCATGGA ATGCCATGGA ATGCCATGGA ATGCCATGGA ATGCCATGGA ATGCCATGGA ATGCCATGGA ATGCCATGGA AAGCAGGCTG AATTCAACTT AAGTGCCAAGAGAA CCTTCACTGA AGTGGTCCCG ACCATATGTA  18 Protein cession #: /	CCTGCTGCTC CCACTTCGAG CCACTTCGAG TTGTGCAAAT CCGCCTGAGA GACATTGAA TCGGATGCAT GTACGCAT GTACGCAT GTACGCAT CCTCTCTCC CTACATCT CCTCTCTCACAT TTCTCACAT TTCTCACAT TTCTCACAT TCCCCTCC GCAAGACAGCA GCACATCAC TCTCCCCTCC GCAAGACACAT TCTCCCCTCC GCTGATGAC TCTCCCTTCC GCTGATGAC TCTCCCTTCC GCAAAACACTAC TCTCCCTTCC GCTGATGAC TCTCCCTTC TTACCCAT TCTCCCTTC TTACCCAT TCATACTAC GCACATTAC TCTCCCTTC TTAC BCACATTAC TCATACTAC TCATACTAC TCATACTAC TTAC TTAC SCACATTAC TTAC	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATTCACGCT GAGAAGTTGTG GTGACTGTGG CGACTCACCA GCCTGCAACC GATGACAACC GATGACAGT ACTGGAAGGA ATACTGAAGT ACTGTCTTTG CCGGATAACC GTCAGCACC GTCAGCTCT TCTTTGACGT ACTTCTCTTT ACTTCTCTTT ACTTCTCTTCT	CTGTGCCCGA TTCAAAAGAT TAGAGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGGATCCCTGA CTACAGCAGC TGGTGGTAGA TCATACTAC TCATTACTAG TCCAGTTCA CCACTATTAC CAATTGGCTT TCCCCAACAT TGTTCCTGTC GGCGCCAGCC CAAGTGGTGGT TCCCCATCAC CCACTCCGA AGGGCCAGGC AGGCCAGCC CTACCGAAAT ATTCCAATGA TGCTTCACCA AGGCCAGGC AGGCCAGCC TTCCGAAAT TTCCAATGA TTTCCAATGA TGCTTCACCA ACAAGAGCGAA TTTCCAATGA TGCTTCACCA TGCTTCACCA TTTCCATTGA TTTCCATTGA TTTCCATTGA TTTCCATTGA TTTCCATTGA TTTCCATTGA TTTCCATTGA TTTAATCCTGA TTTAATCCTGA TTTAGGTTGTT  41   NLFNCKNCAN	GGGCAACTAC AGTCGTCCTG GGACTACTG GGACTACTG GGACTACTT GTTGGACAGG TGTGCCTGTG GGACTACTT GTTTCAGCTT TTGTTCCCTG GAGCTATGTT CCACATGACT CCACATGACT CCACATGACT CTCTTGGATG TCCTGTATC CTTGCTGGAG TAGGCGACAC AGGAAACCTG CCCCGCAGCCCAG CCCCCTGGCC CAGCTATGGT CCGCAACCGT GAGCTTGGAG CTGCCTGCC TGATGACT CCCCCCAACCGT CCCCCCAACC TGATGACTC CCCCCAACCGT CCCCCCCAACC CTACGTCCC CAACATTGTT  51   EAVVQKILDR	120 180 240 300 360 420 600 660 720 780 840 900 1020 1080 1260 1380 1500 1550 1680 1740 1880 1680
<ul><li>50</li><li>55</li><li>60</li><li>65</li><li>70</li><li>75</li></ul>	CCCAGTCCCA AACCTCTTCA GTGCTGTCAA AGAATATCTA ATCACGATGT CTGAACTGG TTGAACAGGT GATCTGCATA TACACGGTTG GATCTGCATA ACAGGTTGC TATTTCTACA AACAGCTACC TCGTTTTGGA AAGGCCACTTG TATGTCTACA AGGAGCCCC CAGGATGGCC GCCCCCTGG CTGGCCACCT ACTGGAGAAA AGTTCGCTTTA GTCGAATGGCC TCGGATGGCC CTCGGATGGCC CTCGGATGGCC TCGGATGGCC TCGGATGGCC TCGGATGGCC TCGGATGGCC TCGGATGGCC TCGGATGGCC TCGGATGGCC TCGGATGGCC TCGGATGAGC TCGGATGGCC TCGGATGGCC TCGGATGGCC TCGGATGGCC TCGGATGGCC TCGGTTGCAAG ATTAAGGAGC ATGGCCCATG CTTGGGTGAAC ATGGCCCATG CTTGGGTGACA TACTGGGTAT Seq ID NO: Protein Ac  I MIRAAVILLL VLSRYDVRLR	TCCCGAAATT ACTGCAAAAAA GATACGATGT TTTATGTCAC TTTTTCATCA CCCTGGACTA AGGATGCTTT GAACGGTGCG AATTCCCTAT AGGACTACTA TCAGGTTCCTA TCAGGTTCCTA TCAGCTCCTA TCAACTACTA TCAACTATCT GAAGAGTCAT TCAACTATCA TCAACTATCA ACTACTCA ACCCCGGA ACCCTGAGCCA ACCCATCGA ATGCCCATGCA ACCCATCGA ACCATCCAA ACCAGCCCAGA ACCATCCAA ACCAGCCCAGA ACCATCCAA ACCAGCCCATGCCA ACCATCCAA ACCAGCCCAACCCAA	CCTGCTGCTC CCACTTCGAG CCACTTCGAG AGACATTGAAA TCGGCATGAGA ATCGGATGCAT CGGCATGAGA TCGGATGCAT CGGCATGAT CGACAGCAG ATTATTCTGG GTTCACTTTC CATACGCCT TTCCTCTGCA CTCACATCTG CCTCACATCTG CCTCACATCTG CCTCACATCTG CCTCACATCTG CCTCACATCTG CCTCACATCTG TTCCCCTCC GGCAGAGCAGA	ATCAGGACCT TTCTCCTCTG GAAGCTGTGG CCGAATTTTG CAGATCTCAG GATCTCACGCT GAGAAGTTGT GTGACTGTGG CGACTCACCC GATGACACC GATGACAATG CCGGGATACC GATGACAATG CCGGGATAAGC TTGTTCTCTCA GCCAGGGTGA ACTCTCTCTCA CCTACAGCC GTCAGCTCTC ACCTCAGAGC ACTATTTTC CACTCAGAGC AGTATTTTC CACCTCAGAGC TCTCTCTCT ACCTCAGAGC TCTCTCTCT ACCTCAGAGC CACAACAATT CCACTCAGAGC CACATGAAG CGCAACACATT CCTCTCGGCCT CCTCTGGCCT  31	CTGTGCCCGA TTCAAAAGAT GAGGTGCCCC AAATGAATAT TAGCATACTA GGGTCCCTGA AGAATCGCGT CTACAGCAGC TCGTGGTAGA TCCACTATTAC CCACTATTAC CCACTATTAC CCACTATTAC GGGCCAGCC AAGTGGTGGT TCCCCAACAT TCCCAACAT TCCCAACAT TCCCAACAT TCCCAACAT ATTCCAATGA TTCCAATGA TTGGGTTGTT  41	GGGCAACTAC AGGCCACC TTTGGACAGG TGTGCCTGTG GGACTACACG TGTGCCTGTG GGACTACTT TTGTTCCCTG GAGCTATCTT TTGTTCACTT CAAGGAGGTG CCACATGACT CAAGGAGGTG TCCTGTATC TTGTTCAATG TTCCTGTATC CTTGCTGATA AACTTCAATG CCTAGCGCAG CCAGCCCAG CCCCCTGCCC CAGCTATGCT CAGCAACCTT CAGCACCAGC CCCCCAACCGT CCCCCAACCGT CCCCCAACCGT CCCCCCAACCGT CCCCCCAACCT CCACCCCAACCT CAACATTGTT CCACCCCAACCCT CAACATTGTT CCACCCCAACCCT CAACATTGTT  51	120 180 240 300 360 420 600 660 720 780 840 900 1020 1080 1260 1380 1500 1550 1680 1740 1880 1680

5	YFYTGSYIRL LILTTIDSHL RRPRRVIARY LATSESLSPL VEAHGHGVTH IKEQFKCDTN	ACNLVVESYG ILKFQVQREV RDKLPNISCI RYQQVVVGNV TSLSGQAPLA DHEDSNESLS STWGLNDDEL PLAFGLFNIV	NSYLVQVYWP KAIDIYILVC QDGLINVEDG TGESLSDLPS SDERHGHGPS MAHGQEKDSS	TVLTTITSWI LFFVFLSLLE VSSLPITPAQ TSEQARHSYG GKPMLHHGEK	SFWMNYDSSA YVYINYLFYS APLASPESLG VRFNGFQADD GVQEAGWDLD	ARVTIGLTSM RGPRRQPRRH SLTSTSEQAQ SIFPTEIRNR DNNDKSDCLA	240 300 360 420 480 540
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15	CGATGTCCGC TAATGAAGAT	11   GCAAATGAAG CTGAGACCGA GAGCTCATGG CCAAGCCCTG	ATTTTGGANN CCCATGGCCA	NATGCTTGCT AGAGAAGGAC	ACTAACAGTA AGTAGCTCAG	CCCGGGGCCT AGTCTGAGGA	60 120 180 240
20	TCCTGACTAC GTTGTTCAAC	GTCCCAAAGG ATTGTAGCGG	TCGACAAGTG CCGAACGATG	GTCCCGGTTC			300
25	-	40 Protein cession #: /	_				
	I       KNCANEAVVQ	11     KILDRVLSRY	21     DVRLRPNFGX	31     MLATNSTRGL	41   NEDELMAHGQ	51   EKDSSSESED	60
30		TEGFSFDLLN					
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35	1	11	21	31	41	51	
		 CACACCTCCC TGAGCCATCC					60 120
40	CGGTGCTGCT	GCTGCTGCTG	CTGCTGCCGC	CACTGCTGCT	GCTGGCGGGG	GCCGTCCCGC	180
40		CCGTGCCGCG					240
		TGCCGACGCC CCAAGGGGAA					300 360
		CTGTGTCCAT					420
45		CATGTTGGCT					480
7.7		GTTTTTCCTG					540 600
	GCCTGAGCTG	CATGAATAAG	GATCACGGCT	GTAGTCACAT	CTGCAAGGAG	GCCCCAAGGG	660
		CTGTGAGTGC TAACCATGGG					720
50		CAGCTGCCAT					780 840
	AGCGAGAGGA	CACTGTCCTG	GAGGTGACAG	AGAGCAACAC	CACATCAGTG	GTGGATGGGG	900
		GAAACGGCGG					960
		TAAGGATACT TGGGAAGACA					1020 1080
55	GTGATCATTT	CTGCAAAAAC	ATCGTGGGCA	GTTTTGACTG	CGGCTGCAAG	AAAGGATTTA	1140
		AGATGAGAAG					1200
		CTGCATCAAC					1260 1320
60		CTGTGTGAAC					1380
60		GAATAAAAAA				ACAAGTGTGT TTCCTCAGAT	1440 1500
						GTAACCTTTA	
		AGGCAAGTGT					1620
65						AACCTTACAT AAGGAAATGT	
00						TCTTGTGACC	
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		CAGGGAGCAG				GACGTGGCTA	1920 1980
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	GCATTTTATG	TCCAAATGGA	ACCTTCCAAA	ATGAGGAAGG	ACAAATGACT	TGTGAACCAT	2100
						ATGTCTGAAT TGCCAGCTCT	
						TGCCAGCTCT	2220 2280
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						CCAGTGGGAA	
						ACTACGACTG GGGGAGCTGG	
00						GCCAACACCG	
80	AGTGTACGTG	GACCATCAAC	CCACCCCCA	AGCGCCGCAT	CCTGATCGTG	GTCCCTGAGA	2640
						ACCTCTTCAT GCCTTCACCT	
						GCTAGAGGGT	

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TCCAGGTCCC ATACGTGACA TATGATGAGG ACTACCAGGA ACTCATTGAA GACATAGTTC
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GGTTGGTGGG ACAGAGCTGT CTTCCTTCTG CATGTCAGCA CAGTCGGGTA TTGCTGCCTC
                                                                                3120
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40	EPTIPRTHLD NRTQQIGVLS NKEIRLIQEE GHSTPKLTPR ALRMTHTLPS KEKARLGQLR	LEEKNVLIQE TSAELRYSVG SHPFESDTEM KESTELRAEE SPAREMDRMG SYHNDARSSL GFMETEAAAQ	SETFRKNLEE SLVDSQSDYR SDIDDDDRET IENRVASVSL VMTLPSDLRK SVSLEPESLG ESLGLGKLGT	SLHDKESLAE TTKVIRRPRR IFSSMDLLSP EGLNLAMVHP HRRKIAVVEE LGSANSSQDS QAEKDRRLKK	EIEKLRSELD GRMGVRRDEP SGHSDAQTLA GTSITASVTA DGREDKATIK LHKAPKKKGI KHELLEEARR	QLKMRTGSLI KVKSLGDHEW MMLQEQLDAI SSLASSSPPS CETSPPPTPR KSSIGRLFGK KGLPFAQWDG	540 600 660 720 780 840
40	EPTIPRTHLD NRTQQIGVLS NKEIRLIQEE GHSTPKLTPR ALRMTHTLPS KEKARLGQLR PTVVAWLELW	LEEKNVLIQE TSAELRYSVG SHPFESDTEM KESTELRAEE SPAREMDRMG SYHNDARSSL GFMETEAAAQ LGMPAWYVAA	SETFRKNLEE SLVDSQSDYR SDIDDDDRET IENRVASVSL VMTLPSDLRK SVSLEPESLG ESLGLGKLGT CRANVKSGAI	SLHDKESLAE TTKVIRRPRR IFSSMDLLSP EGLNLAMVHP HRRKIAVVEE LGSANSSQDS QAEKDRRLKK MSALSDTEIQ	EIEKLRSELD GRMGVRRDEP SGHSDAQTLA GTSITASVTA DGREDKATIK LHKAPKKKGI KHELLEEARR REIGISNPLH	QLKMRTGSLI KVKSLGDHEW MMLQEQLDAI SSLASSSPPS CETSPPPTPR KSSIGRLFGK KGLPFAQWDG RLKLRLAIQE	540 600 660 720 780 840 900 960
40	EPTIPRTHLD NRTQQIGVLS NKEIRLIQEE GHSTPKLTPR ALRMTHTLPS KEKARLGQLR PTVVAWLELW MVSLTSPSAP	LEEKNVLIQE TSAELRYSVG SHPFESDTEM KESTELRAEE SPAREMDRMG SYHNDARSSL GFMETEAAAQ LGMPAWYVAA PTSRTPSGNV	SETFRKNLEE SLVDSQSDYR SDIDDDDRET IENRVASVSL VMTLPSDLRK SVSLEPESLG ESLGLGKLGT CRANVKSGAI WVTHEEMENL	SLHDKESLAE TTKVIRRPRR IFSSMDLLSP EGLNLAMVHP HRRKIAVVEE LGSANSSQDS QAEKDRRLKK MSALSDTEIQ AAPAKTKESE	EIEKLRSELD GRMGVRRDEP SGHSDAQTLA GTSITASVTA DGREDKATIK LHKAPKKKGI KHELLEEARR REIGISNPLH EGSWAQCPVF	QLKMRTGSLI KVKSLGDHEW MMLQEQLDAI SSLASSSPPS CETSPPPTPR KSSIGRLFGK KGLPFAQWDG RLKLRLAIQE LQTLAYGDMN	540 600 660 720 780 840 900
	EPTIPRTHLD NRTQQIGVLS NKEIRLIQEE GHSTPKLTPR ALRMTHTLPS KEKARLGQLR PTVVAWLELW MVSLTSPSAP HEWIGNEWLP	LEEKNVLIQE TSAELRYSVG SHPFESDTEM KESTELRAEE SPAREMDRMG SYHNDARSSL GFMETEAAAQ LGMPAWYVAA PTSRTPSGNV SLGLPQYRSY	SETFRKNLEE SLVDSQSDYR SDIDDDDRET IENRVASVSL VMTLPSDLRK SVSLEPESLG ESLGLGKLGT CRANVKSGA WVTHEEMENL FMECLVDARM	SLHDKESLAE TTKVIRRPRR IFSSMDLLSP EGLNLAMVHP HRRKLAVVEE LGSANSSQDS QAEKDRRLKK MSALSDTEIQ AAPAKTKESE LDHLTKKDLR	EIEKLRSELD GRMGVRRDEP SCHSDAQTLA GTSITASVTA DGREDKATIK LHKAPKKKGI KHELLEEARR REIGISNPLH EGSWAQCPVF VHLKMVDSFH	QLKMRTGSLI KVKSLGDHEW MMLQEQLDAI SSLASSSPPS CETSPPPTPR KSSIGRLFGK KGLPFAQWDG RLKLRLAIQE LQTLAYGDMN RTSLQYGIMC	540 600 660 720 780 840 900 960
40 45	EPTIPRTHLD NRTQQIGVLS NKEIRLIQEE GHSTPKLTPR ALRNTHTLPS KEKARLGQLR PTVVWLELW MVSLTSPSAP HEWIGNEWLP LKRLNYDRKE	LEEKNVLIQE TSAELRYSVG SHPFESDTEM KESTELRAEE SPAREMDRMG SYHNDARSSL GFMETEAAAQ LGMPAWYVAA PTSRTPSGNV SLGLPQYRSY LERRREASOH	SETFRKNLEE SLVDSQSDYR SDLDDDDRFT IEMRVASVSL VMTLPSDLRK SVSLEPESLG ESLGIGKLGT CRANVKSGAI WVTHEEMENL WTHEEMENL EIKDVLVWSN	SLHDKESLAE TTKVIRRPRR IFSSMDLLSP EGLNLAMVHP HRRKIAVVEE LGSANSSQDS QAEKDRLKK MSALSDTEIQ AAPAKTKESE LDHLTKKDLR DRVIRWIQAI	EIEKLRSELD GRMGVRRDER SCHSDAQTLA GTSITASVTA DGREDKATIK LHKAPKKGI KHELLEEARR REIGISNPLH EGSWAQCPVF VHLKMVDSFH GLREYANNIL	QLKMRTGSLI KVKSLGPHEW MMLQEQLDAI SSLASSSPPS CETSPPPTFR KSSIGRLPGK KGLPFAQWDG RLKLRLAIQE LQTLAYGDMN RTSLQYGIMC ESGVHGSLIA	540 600 660 720 780 840 900 960 1020 1080
	EPTIPRTHLD NRTQQIGVLS NKEIRLIQEE GHSTPKLTPR ALRMTHTLPS KEKARLGQLE PTVVWHLELW MVSLTSPSAP HEWIGNEWLP LKRLNYDRKE LDENFDYSSL	LEEKNVLIQE TSAELRYSVG SHPFESDTEM KESTELRAEE SPAREMDRMG SYHNDARSSL GFMETEAAAQ LGMPAWYVAA PTSRTPSGNV SLGLPQYRSY LERRREASQH ALLLQIPTQN	SETFRKNLEE SLVDSQSDYR SDIDDDDRET IEMRVASVSL VMTLPSDLRK SVSLEPESLG CEANVKSGAI WVTHEEMENL FMECLVDARM EIKDVLVWSN TQARQILERE	SLHDKESLAE TTKVIRRPRR IFSSMDLLSP EGLNLAMVHP HRRKLAVVEE LGSANSSQDS QAEKDRRLKK MSALSDTEIQ AAPAKTKESE LDHLTKKDLR	EIEKLRSELD GRMGVRRDEP SCHSDAQTLA GTSITASVTA DGREDKATIK LHKAPKKKGI KHELLEEARR REIGISNPLE EGSWAQCPVF VHLKMVDSFH KRUDSITH KRUDSITH KRUDSIDKN	QLKMRTGSLI KVKSLGDHEW MMLQEQLDAI SSLASSSPPS CETSPPPTRR KSSIGRLFGK KGLPFAQWDG RLKLRLAIQE LQTLAYGDMN RTSLQYGIMC ESGVHGSLIA FRRGSTWRRQ	540 600 660 720 780 840 900 960 1020

It is understood that the examples described above in no way serve to limit the true scope of this invention, but rather are presented for illustrative purposes. All publications, sequences of accession numbers, and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference.

## WHAT IS CLAIMED IS:

1. A method of detecting an androgen-independent prostate cancer cell in a sample from a patient having undergone androgen ablation therapy, the method comprising determining the presence or absence of a nucleic acid comprising a sequence at least 80% identical to a sequence as shown in Tables 1A-4.

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- 2. The method of claim 1, wherein said determining is by hybridizing with a polynucleotide that selectively hybridizes to a sequence at least 95% identical to a sequence as shown in Tables 1A-4.
- 10 3. The method of claim 1, wherein the biological sample:
  - a) is a tissue sample; or
  - b) comprises isolated nucleic acids.
  - 4. The method of claim 3:
- a) wherein the nucleic acids are mRNA; or
  - b) further comprising the step of amplifying nucleic acids before the step of contacting the biological sample with the polynucleotide.
  - 5. The method of claim 2, wherein the polynucleotide:
- a) comprises a sequence as shown in Tables 1A-4;
  - b) is labeled, including a fluorescent label; or
  - c) is immobilized on a solid surface.
- 6. The method according to claim 1, wherein said biological sample is contacted with a plurality of polynucleotides that each selectively hybridizes to a sequence at least 95% identical to a first sequence as shown in Tables 1A-4.
  - 7. The method according to claim 6, wherein said plurality of polynucleotides are immobilized on a solid surface.

8. An isolated polypeptide which is encoded by a nucleic acid molecule having polynucleotide sequence as shown in Tables 1A-4.

9. An antibody that specifically binds a polypeptide of claim 8.

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- 10. The antibody of claim 9:
  - a) further conjugated to an effector component, including a fluorescent label a radioisotope or a cytotoxic chemical; or
  - b) which is an antibody fragment or humanized antibody.

- 11. A method of detecting an androgen-independent prostate cancer cell in a patient having undergone androgen ablation therapy, the method comprising contacting a samp from said patient with an antibody of claim 9.
- 45 12. The method of claim11, wherein:
  - a) the antibody is further conjugated to an effector component, e.g., a fluorescer label; or.
  - b) said sample comprises a cell.
- 13. A method of detecting antibodies specific to androgen-independent prostate can a patient having undergone androgen ablation, the method comprising contacting a biok sample from the patient with a polypeptide encoded by a nucleic acid comprising a sequence from Tables 1A-4.
- 14. A method of inhibiting proliferation of androgen-independent prostate cancer ce a patient having undergone androgen ablation therapy, the method comprising administration to the patient a therapeutically effective amount of a compound that specifically eliminatells expressing an antigen listed in Tables 1A-4.
- 60 15. The method of claim 14, wherein the compound is an antibody.
  - 16. A drug screening assay comprising the steps of:

a) administering a test compound to a mammal having a prostate proliferative condition or a cell isolated therefrom;

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- b) comparing the level of gene expression of a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1A-4 in a treated cell or mammal with the level of gene expression of the polynucleotide in a control cell or mammal, wherein a test compound that modulates the level of expression of the polynucleotide is a candidate for the treatment of prostate cancer.
- 17. The assay of claim 16, wherein:
  - a) the control is a mammal with prostate cancer or a cell therefrom that has not been treated with the test compound; or
  - b) the control is a normal cell or mammal.
- 18. A method for treating a mammal having a prostate proliferative condition or prostate cancer comprising administering a compound identified by the assay of claim 16.
- 80 19. A pharmaceutical composition for treating a mammal having a prostate proliferative condition or prostate cancer, the composition comprising a compound identified by the assay of claim 16 and a physiologically acceptable excipient.
- 20. A method of detecting a prostate cancer associated transcript, the method comprising contacting a biological sample from the patient with a plurality of polynucleotides wherein at least two of said polynucleotides selectively hybridize to a difference sequence at least 80% identical to a sequence as shown in Tables 1A-4.
  - 21. A method of detecting a prostate cancer, the method comprising the steps of:
    - a) providing a biological sample from a patient;
      - b) contacting the biological sample with a first polynucleotide that selectively hybridizes to a sequence at least 80% identical to a first sequence as shown in Tables 1A-4, to determine the level of a prostate cancer-associated transcript in the biological sample; and with a second polynucleotide that selectively

- hybridizes to a second sequence at least 80% identical to a sequence not shown in Tables 1A-4; wherein the expression of said second sequence is not substantially changed in prostate cancer, to determine the level of expression of a control transcript in the biological sample; and
- c) comparing the level of the prostate cancer-associated transcript to a level of the normal tissue associated transcript in the biological sample.
  - 22. A method for quantitation of a prostate cancer-associated transcript in a cell from a patient, the method comprising contacting a biological sample from the patient with a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1A-4.
  - 23. The method of claim 22, wherein:
    - a) the polynucleotide selectively hybridizes to a sequence at least 95% identical to a sequence as shown in Tables 1A-4;
- b) the biological sample is a tissue sample;
  - c) the biological sample comprises isolated nucleic acids;
  - d) the nucleic acids are mRNA;
  - e) further comprising the step of amplifying nucleic acids before the step of contacting the biological sample with the polynucleotide;
  - f) the polynucleotide comprises a sequence as shown in Tables 1A-4;
  - g) the polynucleotide is labeled, including a fluorescent label; or
  - h) the polynucleotide is immobilized on a solid surface.
- 24. A biochip comprising a plurality of polynucleotides that selectively hybridize to a sequence at least 80% identical to a sequence as shown in Tables 1A-4.
  - 25. A method of screening drug candidates comprising:
    - a) providing a cell that expresses an expression profile gene selected from the group consisting of an expression profile gene set forth in Tables 1A-4 or fragment thereof;
    - b) adding a drug candidate to said cell; and

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c) determining the effect of said drug candidate on the expression of said expression profile gene.

130 26. A method according to claim 22 wherein said determining comprises comparing the level of expression in the absence of said drug candidate to the level of expression in the presence of said drug candidate.

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For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

METHODS OF DIAGNOSIS AND TREATMENT OF ANDROGEN-DEPENDENT PROSTATE CANCER, PROSTATE CANCER UNDERGOING ANDROGEN-WITHDRAWAL, AND ANDROGEN-INDEPENDENT PROSTATE **CANCER** 

(57) Abstract: Described herein are genes whose expression are up-regulated or down-regulated in prostate cancer. Also described are such genes whose expression is further up-regulated or down-regulated in drug-resistant prostate cancer cells. Related methods and compositions that can be used for diagnosis and treatment of prostate cancer are disclosed. Also described herein are methods that can be used to identify modulators of prostate cancer.

# INTERNATIONAL SEARCH REPORT

International application No.

PCT/US02/17594

A. CLASSIFICATION OF SUBJECT MATTER						
IPC(7) : C12Q 1/68; G01N 33/00; A61K 31/00; C07K 17/00, 16/00						
	US CL : 435/6, 7.1; 530/350, 387.1; 514/1 According to International Patent Classification (IPC) or to both national classification and IPC					
Minimum do	cumentation searched (classification system followed l	ov classific	ation symbols)			
	35/6, 7.1; 530/350, 387.1; 514/1	,	,			
Documentation	on searched other than minimum documentation to the	extent that	such documents are included i	n the fields searched		
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Electronic da	ata base consulted during the international search (name	e of data h	ase and where practicable sea	rch terms used)		
	ontinuation Sheet	c or data of	ase and, where practicable, sea	ion (cinis asca)		
C. DOC	UMENTS CONSIDERED TO BE RELEVANT					
Category *	Citation of document, with indication, where a	ppropriate,	of the relevant passages	Relevant to claim No.		
X	RAY et al. AIM1, a novel non-lens member of the b	etagamma	-crystallin superfamily, is	1-5,8,22,23,		
	associated with the control of tumorigenicity in huma			670212426		
Y	Acad. Sci. USA. April 1997, Vol. 94, pages 3229-3	5234, espe	cially pages 3230-3231.	6,7,9-21,24-26		
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Further	r documents are listed in the continuation of Box C.		See patent family annex.			
• 5	Special categories of cited documents:	"T"	later document published after the inte	emational filing date or priority		
"A" documen	t defining the general state of the art which is not considered to be		date and not in conflict with the appli-			
	ular relevance	43/7				
"E" earlier ap	oplication or patent published on or after the international filing date	"X"	document of particular relevance; the considered novel or cannot be consider			
"L" documen	t which may throw doubts on priority claim(s) or which is cited to		when the document is taken alone			
establish specified	the publication date of another citation or other special reason (as	"Y"	document of particular relevance; the considered to involve an inventive ste			
			combined with one or more other suc	h documents, such combination		
	t referring to an oral disclosure, use, exhibition or other means		being obvious to a person skilled in the	ne art		
	at published prior to the international filing date but later than the date claimed	"&"	document member of the same patent	family		
	actual completion of the international search	Date of a	mailing of the international sear	rch report		
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	o. (703)305-3230					
Form PCT/IS	A/210 (second sheet) (July 1998)	-V-				

# INTERNATIONAL SEARCH REPORT

International application No.

PCT/US02/17594

Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)				
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:				
1. Claim Nos.: because they relate to subject matter not required to be searched by this Authority, namely:				
Claim Nos.:  because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:				
3. Claim Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).				
Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)				
This International Searching Authority found multiple inventions in this international application, as follows: Please See Continuation Sheet				
<ol> <li>As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.</li> <li>As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.</li> <li>As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.: 1-26 with respect to U83115 of Table 1A</li> </ol>				
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:  Remark on Protest  The additional search fees were accompanied by the applicant's protest.				
No protest accompanied the payment of additional search fees.  Form PCT/ISA/210 (continuation of first sheet(1)) (July 1998)				

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### INTERNATIONAL SEARCH REPORT

### BOX II. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING

Group I, claim(s) 1-7 and 20-24, drawn to methods of detecting a prostate cancer cell or associated transcript.

Group II, claim(s) 8 and 13, drawn to a polypeptide and method of use thereof.

Group III, claim(s) 9-12, drawn to an antibody and method of use thereof.

Group IV, claim(s) 14 and 15, drawn to methods of administering a compound which eliminates cells expressing an antigen.

Group V, claim(s) 16-17 and 25-26, drawn to screening methods for a test compound which modulates expression of particular genes.

Group VI, claim(s) 18-19, drawn to a pharmaceutical compound and method of use thereof in the treatment of a mammal.

This application contains claims directed to more than one species of the generic invention. These species are deemed to lack unity of invention because they are not so linked as to form a single general inventive concept under PCT Rule 13.1.

In order for more than one species to be examined, the appropriate additional examination fees must be paid. The species are as follows:

each of the sequences as shown in Tables 1A-4 represents a single invention.

The claims are deemed to correspond to the species listed above in the following manner: N/A

The following claim(s) are generic: all of claims 1-26.

The inventions listed as Groups I-VI do not relate to a single general inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: the claims of the various groups are not so related as to support unity of invention. Each group has its own special technical feature which is not shared with any other group as follows: I - determining the presence or absence of a nucleic acid; II - polypeptide and method of use thereof; III - antibody and method of use thereof; IV - use of a compound to eliminate cells expressing a specific antigen; V - methods of drug screening based on modulation of gene expression; and VI - pharmaceutical compositions and methods of use thereof in treating a prostate proliferative condition of prostate cancer. Thus, the claims lack the same or corresponding special technical features.

The species listed above do not relate to a single general inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, the species lack the same or corresponding special technical features for the following reasons: the plethora of sequences as shown in Tables 1A-4 have no relation whatsoever with one another and thus clearly do not share any special technical features. As each sequence is different from and independent of the others, the particular sequence and biological function of each such sequence constitutes its own special technical features.

Continuation of B. FIELDS SEARCHED Item 3: USPAT, DERWENT WPI, BIOSIS, MEDLINE

search terms: AIM1, accession # U83115

Form PCT/ISA/210 (second sheet) (July 1998)